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(54) Title: MODIFIED GENE-SILENCING RNA AND USES THEREOF

(57) Abstract: Methods and means for efficiently downregulating the expression of any gene of interest in eukaryotic cells and organisms are provided. To this end, the invention provides modified antisense and sense RNA molecules, chimeric genes encoding such modified antisense or sense RNA molecules and eukaryotic organisms such as plants, animals or fungi, yeast or molds comprising the modified antisense and/or sense RNA molecules or the encoding chimeric genes.

### Modified gene-silencing RNA and uses thereof

#### Field of the invention

The present invention relates to methods for efficienty downregulating the expression of any gene of interest in eukaryotic cells and organisms. To this end, the invention provides modified antisense and sense RNA molecules, chimeric genes encoding such modified antisense or sense RNA molecules and eukaryotic organisms such as plants, animals or fungi, yeast or molds comprising the modified antisense and/or sense RNA molecules or the encoding chimeric genes.

#### Background art

- Recently, it has been shown that introduction of double stranded RNA (dsRNA) also called interfering RNA (RNAi), or hairpin RNA is an effective trigger for the induction of gene-silencing in a large number of eukaryotic organisms, including animals, fungi or plants.
- Both the qualitative level of dsRNA mediated gene silencing (level of genesilencing within an organism) and the quantitative level (number of organisms showing a significant level of gene-silencing within a population) have proven superior to the more conventional antisense RNA or sense RNA mediated gene silencing methods.

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For practical purposes, the production of antisense RNA molecules and chimeric genes encoding such antisense RNA is more straightforward than the production of dsRNA molecules or the encoding genes. Indeed, the chimeric nucleic dsRNA molecules or the encoding genes contain large, more or less perfect inverted repeat structures, and such structures tend to hamper the intact maintenance of these nucleic acids in the intermediate prokaryotic

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cloning hosts. The methods and means as hereinafter described to increase the efficiency of antisense-RNA mediated gene silencing provide a solution to this problem as described in the different embodiments and claims.

US 5,190,131 and EP 0 467 349 A1 describe methods and means to regulate or inhibit gene expression in a cell by incorporating into or associating with the genetic material of the cell a non-native nucleic acid sequence. Said sequence is transcribed to produce an mRNA which is complementary to and capable of binding to the mRNA produced by the genetic material of that cell.

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EP 0 223 399 A1 describes methods to effect useful somatic changes in plants by causing the transcription in the plant cells of negative RNA strands which are substantially complementary to a target RNA strand. The target RNA strand can be a mRNA transcript created in gene expression, a viral RNA, or other RNA present in the plant cells. The negative RNA strand is complementary to at least a portion of the target RNA strand to inhibit its activity *in vivo*.

EP 0 240 208 describes a method to regulate expression of genes encoded for in plant cell genomes, achieved by integration of a gene under the transcriptional control of a promoter which is functional in the host. In this method, the transcribed strand of DNA is complementary to the strand of DNA that is transcribed from the endogenous gene(s) one wishes to regulate.

WO95/15394 and US 5908779 describe a method and construct for regulating gene expression through inhibition by nuclear antisense RNA in (mouse) cells. The construct comprises a promoter, antisense sequences, and a cis-or transribozyme which generates 3'-ends independently of the polyadenylation machinery and thereby inhibits the transport of the RNA molecule to the cytoplasm.

WO98/05770 discloses antisense RNA with special secondary structures such as  $(GC)_n$ -palindrome- $(GC)_n$  or  $(AT)_n$ -palindrome- $(AT)_n$  or  $(CG)_n$ -palindrome- $(CG)_n$  and the like.

WO 01/12824 discloses methods and means for reducing the phenotypic expression of a nucleic acid of interest in eukaryotic cells, particularly in plant cells, by providing aberrant, preferably unpolyadenylated, target-specific RNA to the nucleus of the host cell. Preferably, the unpolyadenylated target-specific RNA is provided by transcription of a chimeric gene comprising a promoter, a DNA region encoding the target-specific RNA, a self-splicing ribozyme and a DNA region involved in 3' end formation and polyadenylation.

WO 02/10365 provides a method for gene suppression in eukaryotes by transformation with a recombinant construct containing a promoter, at least one antisense and/or sense nucleotide sequence for the gene(s) to be suppressed, wherein the nucleus-to-cytoplasm transport of the transcription products of the construct is inhibited. In one embodiment, nucleus-to-cytoplasm transport is inhibited by the absence of a normal 3' UTR. The construct can optionally include at least one self-cleaving ribozyme. The construct can also optionally include sense and/or antisense sequences to multiple genes that are to be simultaneously down-regulated using a single promoter. Also disclosed are vectors, plants, animals, seeds, gametes, and embryos containing the recombinant constructs.

Zhao et al., J. Gen. Virology, 82, 1491-1497 (2001) described the use of a vector based on Potato Virus X in a whole plant assay to demonstrate nuclear targeting of Potato spindle tuber viroid (PSTVd).

WO 02/00894 relates to gene silencing methods wherein the nucleic acid constructs comprise within the transcribed region a DNA sequence which consists of a stretch of T bases in the transcribed strand.

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WO 02/00904 relates to gene silencing methods wherein nucleic acid constructs comprise (or encode) homology to at least one target mRNA expressed by a host, and in the proximity thereto, two complementary RNA regions which are unrelated to any endogenous RNA in the host.

#### Summary of the invention

- In one embodiment of the invention a method for down regulating the expression of a target gene in cells of a eukaryotic organisms is provided, comprising the steps of
  - a) providing the cells of the eukaryotic organism with a chimeric RNA molecule wherein the chimeric RNA molecule comprises
    - i) one target-gene specific region or multiple target-gene specific regions comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequence identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene; operably linked to
    - ii) a largely double stranded RNA region comprising a nuclear localization signal from a viroid of the Potato spindle tuber viroid (PSTVd)-type such as Potato Spindle tuber viroid, Citrus viroid species III, Citrus viroid species IV, Hop latent viroid, Australian grapevine viroid, Tomato planta macho viroid, Coconut tinangaja viroid, Tomato apical stunt viroid, Coconut cadang-cadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid and Citrus bent leaf viroid or a the largely double stranded RNA region or a largely double stranded RNA region comprising at least 35 repeats of the trinucleotides CUG, CAG, GAC or GUC such as between 44 and 2000 repeats of these trinucleotide; and

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b) identifying those eukaryotic organisms wherein the expression of the target gene is down regulated.

The chimeric RNA molecule may comprise an intron sequence. The viroids may have a genomic nucleotide sequence selected from the group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8. The eukaryotic organism may be a plant including a plant selected from Arabidopsis, alfalfa, barley, bean, corn, cotton, flax, pea, rape, rice, rye, safflower, sorghum, soybean, sunflower, tobacco, wheat, asparagus, beet, broccoli, cabbage, carrot, cauliflower, celery, cucmber, eggplant, lettuce, onion, oilseed rape, pepper, potato, pumpkin, radish, spinach, squash, tomato, zucchini, almond, apple, apricot, banana, blackberry, b lueberry, cacao, cherry, coconut, cranberry, date, grape, grapefruit, guava, kiwi, lemon, lime, mango. melon, nectarine, orange, papaya, passion fruit, peach, peanut, pear, pineapple, pistachio, plum, raspberry, strawberry, tangerine, walnut and watermelon. The eukaryotic organism may also be a fungus, yeast or mold or an animal such as a human, mammal, fish, cattle, goat, pig, sheep, rodent, hamster, mouse, rat, guinea pig, rabbit, primate, nematode, shellfish, prawn, crab, lobster, insect, fruit fly, Coleapteran insect, Dipteran insect, Lepidopteran insect and Homeopteran insect.

It is an object of the invention to provide a chimeric RNA molecule for down-regulating the expression of a target gene in a cell of a eukaryotic organisms, comprising one target-gene specific region or multiple target-gene specific regions a target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene; operably linked to a largely double stranded RNA region comprising a nuclear localization signal from a viroid of the Potato spindle tuber viroid (PSTVd)-type such as Potato Spindle tuber viroid, Citrus viroid species III, Citrus viroid species IV, Hop latent viroid, Australian

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grapevine viroid, Tomato planta macho viroid, Coconut tinangaja viroid, Tomato apical stunt viroid, Coconut cadang-cadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid and Citrus bent leaf viroid or a the largely double stranded RNA region or a largely double stranded RNA region comprising at least 35 repeats of the trinucleotide CUG, CAG, GAC OR GUC such as between 44 and 2000 repeats of the trinucleotide CUG, CAG, GAC OR GUC wherein the chimeric RNA molecule, when provided to cells of the eukaryotic organism down-regulates the expression of the target gene.

It is another object of the invention to provide a chimeric DNA molecule for reduction of the expression of a target gene in a cell of a eukaryotic organism, comprising

- a) a promoter or promoter region capable of being recognized by RNA polymerases in the cells of the eukaryotic organism; operably linked to
- b) a DNA region, which when transcribed yields an RNA molecule, the RNA molecule comprising
  - i) one target-gene specific region or multiple target-gene specific regions comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene; operably linked to
  - ii) a largely double stranded RNA region comprising a nuclear localization signal from a viroid of the Potato spindle tuber viroid (PSTVd)-type such as Potato Spindle tuber viroid, Citrus viroid species III, Citrus viroid species IV, Hop latent viroid, Australian grapevine viroid, Tomato planta macho viroid, Coconut tinangaja viroid, Tomato apical stunt viroid, Coconut cadang-cadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid and Citrus bent leaf viroid or a the largely double stranded RNA region or a largely double stranded RNA region comprising at least 35 repeats of the trinucleotide CUG, CAG, GAC OR GUC such as between 44

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and 2000 repeats of the trinucleotide CUG, CAG, GAC OR GUC; and optionally

iii) further comprising a transcription termination and polyadenylation signal operably linked to the DNA region encoding the RNA molecule.

wherein the chimeric DNA molecule, when provided to cells of the eukaryotic organism reduces the expression of the target gene.

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Depending on the eukaryotic host organism, the promoter or promoter region may a promoter which functions in animals, a promoter which functions in yeast, fungi or molds or a plant-expressible promoter. The promoter may also be a promoter or promoter region recognized by a single subunit bacteriophage RNA polymerase.

The invention also provides cells from a eukaryotic organism comprising a chimeric DNA or RNA molecules according to the invention, as well as non-human eukaryotic organisms, comprising in their cells a chimeric DNA or RNA molecule according to the invention.

It is yet another object of the invention to provide the use of a chimeric RNA or DNA molecule according to the invention for reduction of the expression of a target gene in a cell of a eukaryotic organism.

The invention also provides a method for making a transgenic eukaryotic organism wherein expression of a target gene in cells of the organism is reduced, the method comprising the steps of :

- a) providing a chimeric DNA molecule according to the invention to a cell or cells of the organism to make a transgenic cell or cells; and
- b) growing or regenerating a transgenic eukaryotic organism from the transgenic cell or cells.

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The invention also provides a method for down regulating the expression of a target gene in cells of a eukaryotic organisms, comprising the steps of

- a) providing the cells of the eukaryotic organism with a first and second chimeric RNA molecule, wherein
  - i) the first chimeric RNA molecule comprises an antisense target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene;
  - ii) the second chimeric RNA molecule comprises a sense target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity to the complement of the first chimeric RNA molecule;
  - iii) the first and second chimeric RNA are capable of basepairing at least between the 19 consecutive nucleotides of the first chimeric RNA and the 19 consecutive nucleotides of the second chimeric RNA; and
  - iv) wherein either the first or the second chimeric RNA molecule comprises a largely double stranded RNA region operably linked to the antisense target-specific RNA region or to the sense targetspecific RNA region; and
- b) identifying those eukaryotic organisms wherein the expression of the target gene is down regulated.

Both the first and second chimeric RNA molecule may comprise a largely double stranded region.

It is another object of the invention to provide a cell from a eukaryotic organism, (as well as non-human eukaryotic organisms comprising such cells), comprising a first and second chimeric RNA molecule,

i) the first chimeric RNA molecule comprising an antisense target-gene specific RNA region comprising a nucleotide sequence of at least

about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene;

- ii) the second chimeric RNA molecule comprising a sense target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity to the complement of the first chimeric RNA molecule;
- iii) the first and second chimeric RNA being capable of basepairing at least between the 19 consecutive nucleotides of the first chimeric RNA and the 19 consecutive nucleotides of the second chimeric RNA; and

wherein either the first or the second chimeric RNA molecule comprises a largely double stranded RNA region operably linked to the antisense target-specific RNA region or to the sense target-specific RNA region.

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The invention further provides chimeric sense RNA molecules or chimeric DNA molecules encoding such chimeric sense RNA molecules for reduction of expression of a target gene in a cell of a eukaryotic organism in cooperation with a chimeric antisense RNA molecule, wherein the chimeric sense RNA molecule comprises a sense target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity to the nucleotide of said target gene; operably linked to a largely double stranded RNA region.

#### 25 Brief description of the figures.

Figure 1: Schematic representation of the secondary structure predicted using Mfold software for different viroids of the PSTVd-type. A. Potato spindle tuber viroid; B. Australian grapevine viroid; C. Coconut tinangaja viroid; D. Tomato planta macho viroid; E. Hop latent viroid of thermomutant T229; F. Tomato apical stunt viroid.

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Figure 2: schematic representation of the various chimeric gene constructs used in the examples 1 to 3 of this application. 35S-P: CaMV 35S promoter; Pdk intron: Flaveria trinervia pyruvate orthophosphate dikinase 2 intron 2; cEIN2: cDNA copy of the EIN2 gene from Arabidopsis (gene required for sensitivity to ehylene; Alonso et al. 1999 Science 284, 2148-2152) the orientation of this region with respect to the promoter is indicated by the arrow; gEIN2: genomic copy of the EIN2 gene from Arabidopsis; PSTVd: cDNA copy of the genome of potato spindle tuber viroid; PSTVd\*: partial sequence from PSTVd from nucleotide 16 to nucleotide 355, cloned in inverse orientation with regard to the intact copy of PSTVd; OCS 3': 3' region of the octopine synthase gene from Agrobacterium tumefaciens.

Figure 3: Phenotype of EIN2-silenced plants when germinating on 1-aminocyclopropane-1-carboxylic acid (ACC). A. In the dark; B. under light conditions. Wt: wild-type plants.

Figure 4: schematic representation of the various chimeric gene constructs used in Example 4. CMV promoter: cytomegolovirus promoter; SV40 poly(A): transcription termination and polyadenylation region from SV40; PSTVd: potato spindle tuber viroid sequence; CUGrep: sequence comprising 60 repeats of the CUG sequence; humGFP: humanized green fluorescent protein coding region (adapted to the codon usage of human genes; the orientation of this region with respect to the promoter is indicated by the arrow);

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Fig 5: Schematic representation of the predicted secondary structure of pSTVd in pMBW491 (A ;adopting almost the wild type configuration) and in pMBW489, where a 10 nucleotide deletion results in a structure different from the wild type configuration.

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#### Detailed description of the different embodiments.

The currently described method and means for obtaining enhanced antisense RNA -mediated down regulation of gene expression are based upon the unexpected observation that operably linking the target gene-specific RNA sequence to a largely double stranded RNA region, such as an RNA region comprising the nucleotide sequence of a Potato spindle tuber viroid genome, which in turn comprises a nuclear localization signal for the RNA in which it is embedded, when introduced into cells of a host organism, such as a plant cell, increased both the number of lines wherein gene expression of the target gene was down-regulated, as well as the number of lines wherein gene expression of the target gene was significantly downregulated or even abolished.

Thus, in one embodiment of the invention, a method is provided for down regulating the expression of a target gene in cells of a eukaryotic organisms, comprising the steps of

- a) providing the cells of the eukaryotic organism with a chimeric RNA molecule wherein the RNA molecule comprises
  - i) a target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene ( the « antisense RNA »); operably linked to
  - ii) a largely double stranded RNA region; and
- b) identifying those eukaryotic organisms wherein the expression of the target gene is down regulated.
- "Chimeric gene " or "chimeric nucleic acid " as used herein, refers any gene or any nucleic acid, which is not normally found in a particular eukaryotic species or, alternatively, any gene in which the promoter is not associated in

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nature with part or all of the transcribed DNA region or with at least one other regulatory region of the gene.

As used herein, "antisense RNA" refers to RNA molecules which comprise a nucleotide sequence that is largely complementary to part of the nucleotide sequence of the biologically active RNA, usually but not exclusively mRNA, which is transcribed from the target gene.

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The expression « target gene » is used herein to refer to any nucleic acid which is present in the eukaryotic cell and that is transcribed into a biologically active RNA. The target gene may be an endogenous gene, it may be a transgene that was introduced through human intervention in the ancestors of the eukaryotic cell, or it may be a gene introduced into the genome of the cell by infectious organisms such as e.g. *Agrobacterium* strains or retroviruses. The target gene may also be of viral origin. Furthermore, the stretch of at least 19 nucleotides may be selected from the promoter region, the 5'UTR, the coding region, or the 3'UTR.

"Gene expression" or "expression of a nucleic acid" is used herein to refer to the process wherein a gene or nucleic acid, when introduced in a suitable host cell, can be transcribed (or replicated) to yield an RNA, and/or translated to yield a polypeptide or protein in that host cell.

As used herein, "downregulation of gene expression" refers to the comparison of the expression of the target gene or nucleic acid of interest in the eukaryotic cell in the presence of the RNA or chimeric genes of the invention, to the expression of target gene or the nucleic acid of interest in the absence of the RNA or chimeric genes of the invention. The expression of the target gene in the presence of the chimeric RNA of the invention should thus be lower than the expression in absence thereof, such as be only about 50% or 25% or about 10% or about 5% of the phenotypic expression in absence of the chimeric RNA. For a number of applications, the expression should be completely

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inhibited for all practical purposes by the presence of the chimeric RNA or the chimeric gene encoding such an RNA.

As used herein "comprising" is to be interpreted as specifying the presence of the stated features, integers, steps or components as referred to, but does not preclude the presence or addition of one or more features, integers, steps or components, or groups thereof. Thus, e.g., a nucleic acid or protein comprising a sequence of nucleotides or amino acids, may comprise more nucleotides or amino acids than the actually cited ones, i.e., be embedded in a larger nucleic acid or protein. A chimeric gene comprising a DNA region which is functionally or structurally defined, may comprise additional DNA regions etc.

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It will thus be clear that the minimum nucleotide sequence of the antisense RNA of about 19 nt of the target-gene specific RNA region may be comprised within a larger RNA molecule, varying in size from 19 nt to a length equal to the size of the target gene with a varying overall degree of sequence identity.

For the purpose of this invention, the "sequence identity" of two related nucleotide or amino acid sequences, expressed as a percentage, refers to the number of positions in the two optimally aligned sequences which have identical residues (x100) divided by the number of positions compared. A gap, i.e., a position in an alignment where a residue is present in one sequence but not in the other is regarded as a position with non-identical residues. The alignment of the two sequences is performed by the Needleman and Wunsch algorithm (Needleman and Wunsch 1970) The computer-assisted sequence alignment above, can be conveniently performed using standard software program such as GAP which is part of the Wisconsin Package Version 10.1 (Genetics Computer Group, Madision, Wisconsin, USA) using the default scoring matrix with a gap creation penalty of 50 and a gap extension penalty of 3. Sequences are indicated as "essentially similar" when such sequence have a sequence identity of at least about 75%, particularly at least about 80 %, more

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particularly at least about 85%, quite particularly about 90%, especially about 95%, more especially about 100%, quite especially are identical. It is clear than when RNA sequences are the to be essentially similar or have a certain degree of sequence identity with DNA sequences, thymine (T) in the DNA sequence is considered equal to uracil (U) in the RNA sequence. Thus when it is stated in this application that a sequence of 19 consecutive nucleotides has a 94% sequence identity to a sequence of 19 nucleotides, this means that at least 18 of the 19 nucleotides of the first sequence are identical to 18 of the 19 nucleotides of the second sequence.

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The mentioned antisense nucleotide regions may thus be about 21nt, 50 nt, 100nt, 200 nt, 300nt, 500nt, 1000 nt, 2000 nt or even about 5000 nt or larger in length, each having an overall sequence identity of about 40 % or 50% or 60 % or 70% or 80% or 90 % or 100%. The longer the sequence, the less stringent the requirement for the overall sequence identity is.

Furthermore, multiple sequences with sequence identity to the complement of the nucleotide sequence of a target gene (multiple target-gene specific RNA regions) may be present within one RNA molecule. Also, multiple sequences with sequence identity to the complement of the nucleotide sequences of several target genes may be present within one RNA molecule.

"Target-gene specific" is not to be interpreted in the sense that the chimeric nucleic acids according to the invention can only be used for down-regulation of that specific target gene. Indeed, when sufficient homology exists between the target gene specific RNA region and another gene, or when other genes share the same stretch of 19 nucleotides (such as genes belonging to a so-called gene-family) expression of those other genes may also be down-regulated.

As used herein, a « largely double stranded RNA region » refers to an RNA molecule which is capable of folding into a rod-like structure by internal base-

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pairing and wherein the resulting rod-like structure does not comprise any stretch of 19 consecutive nucleotides having 94% sequence identity to the complement of another stretch of 19 other consecutive nucleotides within that RNA molecule, which are capable of forming a double stranded region when the RNA molecule folds into a rod-like structure. In other words, the largely double stranded RNA region upon folding does not contain a double stranded RNA regions of at least 19 bp with at most one mismatch in those 19 bp, at least not in the energicatically most favourable rod-like confirmation. Non-limiting examples of such structures are represented in figure 1.

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Although not intending to limit the invention to a specific mode of action, it is thought that such largely double stranded RNA regions are involved in the nuclear localization of the antisense RNA molecules with which they are associated. As a consequence thereof, the concentration of the antisense RNA molecules in the nucleus may be increased, allowing a more efficient formation of the formation of sequence specific dsRNA formation by base pairing with the sense RNA corresponding to the antisense RNA.

As used herein, the term « Capable of folding into a rod-like structure » with regard to an RNA molecule refers to a secondary structure which the RNA molecule will preferably adapt by internal basepairing and which has the overall appearance of a long rod. The rod-like structure may comprise branches or bulges (where non-matching nucleotides bulge out from the overall structure) and may be part of a larger secondary structure (which may or may not be rod-like). Examples of RNA molecules capable of folding into a rod-like structure are represented in Figure 1.

The specific secondary structure adapted will be determined by the free energy of the RNA molecule, and can be predicted for different situations using appropriate software such as FOLDRNA (Zuker and Stiegler, 1981) or the

MFOLD structure prediction package of GCG (Genetics Computing Group; Zuker 1989, Science 244, 48-52).

In one embodiment of the invention, the largely double stranded RNA region operably linked to the antisense RNA molecule is a nuclear localization signal from a viroid of the PSTVd type, such as PSTVd (Potato spindle tuber viroid), capable of replicating in the nucleus of the host cell or host plant cell.

In one embodiment of the invention, the largely double stranded RNA region comprises the full length sequence of PSTVd strain RG1, which can conveniently be obtained by amplification from a cDNA copy of the RNA genome of the viroid using oligonucleotide primers with the nucleotide sequence

5'-cgcagatctcggaactaaactcgtggttc-3' [SEQ ID N°1] and 5'gcgagatctaggaaccaactgcggttc-3'[SEQ ID N°2]), such as the nucleotide sequence represented in SEQ ID N°3.

It is understood that for incorporation in an RNA molecule, an additional step is required to convert the DNA molecule in the corresponding RNA molecule. Such a conversion may be achieved by transcription, e.g. in vitro transcription using a single subunit bacteriophage RNA polymerase.

It is also clear than when RNA sequences are said to be represented in an entry in the Sequence Listing or to be essentially similar or have a certain degree of sequence identity with DNA sequences represented in the Sequence Listing, reference is made to RNA sequences corresponding to the sequences in the entries, except that thymine (T) in the DNA sequence is replaced by uracil (U) in the RNA sequence. Whether the reference is to RNA or DNA sequence will be immediately apparent by the context.

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Similar largely double stranded RNA structures are also found within the genomes of other nuclear-replicating viroids of the PSTVd type (or group B according to the classification by Bussière et al. 1996) and these RNA sequences may be used to similar effect. Other nuclear-replicating viroids of the PSTVd group include Citrus viroid species III, Citrus viroid species IV, Coleus viroid, Hop latent viroid (SEQ ID N° 7), Australian grapevine viroid (SEQ ID N° 4), Tomato planta macho viroid (SEQ ID N° 6), Coconut tinangaja viroid (SEQ ID N° 5), Tomato apical stunt viroid (SEQ ID N° 8), Coconut cadang-cadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid or Citrus bent leaf viroid. These viroids are also characterized by the absence of self-splicing activity which becomes apparent by the absence of catalytic motifs such as the hammerhead motif (Busière et al. Nuc. Acids Res. 24, 1793-1798, 1996). The longest stretch of perfect dsRNA structures among all the PSTVd-type of viroids is 11 base pairs in size. The mismatches are usually quite evenly distributed.

Nucleotide sequences for these viroids have been compiled in a database accesible via the worldwide web (http://www.callisto.si.usherb.ca/~jpperra or http://nt.ars-grin.gov/subviral/) and include the following:

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Potato spindle tuber viroid (PSTVd) [PSTVd.1 (Accession numbers: J02287(gb), M16826(gb), V01465(embl); 333351(gi), 333352(gi) and 62283(gi)); PSTVd.2 (Accession numbers: M38345(gb), 333354(gi)); PSTVd.3 (Accession numbers: M36163(gb), 333356(gi)); PSTVd.4 (Accession numbers: M14814(gb), 333357(gi)); PSTVd.5 (strain: S.commersonii) (Accession numbers: M25199(gb), 333355(gi)); PSTVd.6 (strain: tomato cv. Rutgers,isolate: KF440-2) (Accession numbers: X58388(embl), 61366(gi)); PSTVd.7 (mild strain KF6-M ) (Accession number: M88681(gb), 333358(gi)); PSTVd.8 (strain Burdock) (Accession numbers: M88678(gb), 333360(gi)); PSTVd.9 (strain Wisconsin (WB)) (Accession numbers: M88677(gb), 333359(gi)); PSTVd.10 (strain PSTVd-N(Naaldwijk)) (Accession numbers:

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X17268(embl), 60649(gi)) ;PSTVd.11 (mild strain variant A, WA-M isolate) (Accession numbers: X52036(embl), 61365(gi)); PSTVd.12 (mild strain, F-M isolate) (Accession X52037(embl), numbers: 61367(gi)); (intermediate-severe strain, F-IS isolate) (Accession numbers: X52039(embl), 61369(gi)); PSTVd.14 (severe-lethal strain, F-SL isolate) (Accession numbers: X52038(embl), 61368(gi)); PSTVd.15 (intermediate-severe strain, F88-IS isolate) as published in Herold, T et al., Plant Mol. Biol. 19, 329-333 (1992); PSTVd.16 (variant F88 or S88)(Accession numbers: X52040(embl), 61370(gi)); PSTVd.17 (individual isolate kf 5) (Accession numbers: M93685(gb), 333353(gi)); PSTVd.18 (isolate KF5) (Accession numbers: S54933(gb), 265593(gi)); PSTVd.19 (strain S-XII, variety s27) (Accession numbers: X76845(embl), 639994(gi)); PSTVd.20 (strain S-XIII, variety s23) (Accession numbers: X76846(embl), 639993(gi)); PSTVd.21 (strain M(mild)) (Accession numbers: X76844(embl), 639992(gi)); PSTVd.22 (strain I-818, variety I4) (Accession numbers: X76848(embl), 639991(gi)); PSTVd.23 (strain I-818, variety I3) (Accession numbers: X76847(embl), 639990(gi)); PSTVd.24 (strain PSTVd-341) (Accession numbers: Z34272(embl), 499191(gi)); PSTVd.25 (strain QF B) (Accession numbers: U23060(gb), 755586(gi)) PSTVd.26 (strain QF A) (Accession numbers: U23059(gb), 755585(gi)); PSTVd.27 (strain RG 1) (Accession numbers: U23058(gb), 755584(gi)); PSTVd.28 (Accession numbers: U51895(gb), 1272375(gi)); PSTVd.29(Potato spindle tuber viroid) (Accession numbers: X97387(embl), 1769438(gi)); PSTVd.30 (strain S27-VI-24) (Accession numbers: Y09382(emb), 2154945(gi)); PSTVd.31 (strain S27-VI-19) (Accession numbers: Y09383(emb), 2154944(gi)); PSTVd.32 (strain SXIII) (Accession numbers: Y08852(emb), 2154943(gi)); PSTVd.33 (strain S27-I-8) (Accession numbers: Y09381(emb), 2154942(gi)); PSTVd.34 (strain PSTV M-VI-15) (Accession numbers: Y09577(emb), 2154941(gi)); PSTVd.35 (strain PSTV M-I-40) (Accession numbers: Y09576(emb), 2154940(gi)); PSTVd.36 (strain PSTV M-I-17) (Accession numbers: Y09575(emb), 2154939(gi)); PSTVd.37 (strain PSTV M-I-10) (Accession Y09574(emb), 2154938(gi)); PSTVd.38 (variant I4-I-42) (Accession numbers:

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Y09889(emb), 2154937(gi)); PSTVd.39 (variant PSTVd I2-VI-27) (Accession numbers: Y09888(emb), 2154936(gi)); PSTVd.40 (variant PSTVd I2-VI-25) (Accession numbers: Y09887(emb), 2154935(gi)); PSTVd.41 (variant PSTVd I2-VI-16) (Accession numbers: Y09886(emb), 2154934(gi)); PSTVd.42 (variant PSTVd I4-I-10) (Accession numbers: Y09890(emb), 2154933(gi)); PSTVd.43 (variant PSTVd I2-I-14) (Accession numbers: Y09891(emb), 2154932(gi)); PSTVd.44 (isolate KF7) (Accession numbers: AJ007489(emb), 3367737(gi)); PSTVd.45 (Accession numbers: AF369530, 14133876(gi)];

Group III citrus viroid (CVd-III) [CVd-III.1 (Accession numbers: S76452(gb), 913161(gi)); CVd-III.2 (Australia New South Wales isolate) (Accession numbers: S75465(gb) and S76454(gb), 914078(gi) and 913162(gi)); CVd-III.3 (Accession numbers: AF123879, GI:7105753); CVd-III.4 (Accession numbers: AF123878, GI:7105752) CVd-III.5 (Accession numbers: AF123877, GI:7105751); CVd-III.6 (Accession numbers: AF123876, GI:7105750); CVd-III.7 (Accession numbers: AF123875, GI:7105749); CVd-III.8 (Accession numbers: GI:7105748); CVd-III.9 (Accession numbers: AF123873, AF123874. GI:7105747); CVd-III.10 (Accession numbers: AF123872, GI:7105746); CVd-III.11 (Accession numbers: AF123871, GI:7105745); CVd-III.12 (Accession numbers: AF123870, GI:7105744); CVd-III.13 (Accession numbers: AF123869, GI:7105743); CVd-III.14 (Accession numbers: AF123868, GI:7105742); CVd-III.15 (Accession numbers: AF123867, GI:7105741); CVd-III.16 (Accession numbers: AF123866, GI:7105740); CVd-III.17 (Accession numbers: AF123865, GI:7105739); CVd-III.18 (Accession numbers: AF123864, GI:7105738) CVd-III.19 (Accession numbers: AF123863, GI:7105737); CVd-III.20 (Accession numbers: AF123860, GI:7105736); CVd-III.21 (Accession numbers: AF123859, GI:7105735); CVd-III.22 (Accession numbers: AF123858, GI:7105734); CVd-III.23 (Accession numbers: AB054619, GI:13537479); CVd-III.24 (Accession GI:13537480); CVd-III.25 (Accession numbers: AB054620, AB054621, GI:13537481); CVd-III.26 (Accession numbers: AB054622, GI:13537482); CVd-III.27 (Accession numbers: AB054623, GI:13537483); CVd-III.28 (Accession numbers: AB054624, GI:13537484); CVd-III.29

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(Accession numbers: AB054625, GI:13537485); CVd-III.30 (Accession numbers: AB054626, GI:13537486); CVd-III.31 (Accession numbers: AB054627, GI:13537487); CVd-III.32 (Accession numbers: AB054628, GI:13537488); CVd-III.33 (Accession numbers: AB054629, GI:13537489); CVd-III.34 (Accession numbers: AB054630, GI:13537490); CVd-III.35 (Accession numbers: AB054631, GI:13537491); CVd-III.36 (Accession numbers: AB054632, GI:13537492); CVd-III.37 (Accession numbers: AF416552, GI:15811643); CVd-III.38 (Accession numbers: AF416553, GI:15811644); CVd-III.39 (Accession numbers: AF416374, GI:15788948); CVd-III.40 (Accession number: AF434680)];

Citrus viroid IV (CVdIV) [CVdIV.1 (Accession numbers: X14638(embl), 59042(gi))]

Coleus blumei-1 viroid (CbVd-1) [CbVd.1 (Coleus blumei viroid 1 (CbVd 1),strain cultivar Bienvenue, german isolate) (Accession numbers: X52960(embl), 58844(gi)); CbVd.2 (Coleus yellow viroid (CYVd), Brazilian isolate) (Accession numbers: X69293(embl), 59053(gi)); CbVd.3 (Coleus blumei viroid 1-RG stem-loop RNA.) (Accession numbers: X95291(embl), 1770104(gi)); CbVd.4 (Coleus blumei viroid 1-RL RNA) (Accession numbers: X95366(embl), 1770106(gi))]

Coleus blumei-2 viroid (CbVd-2) [CbVd.1 (Coleus blumei viroid 2-RL RNA) (Accession numbers: X95365(embl), 1770107(gi)); CbVd.2 (Coleus blumei viroid CbVd 4-1 RNA) (Accession mumbers: X97202(embl), 1770109(gi))] Coleus blumei-3 viroid (CbVd-3) [CbVd.1 (Coleus blumei viroid 3-RL) (Accession mumbers: X95364(embl), 1770108(gi)); CbVd.2 (Coleus blumei viroid 8 from the Coleus blumei cultivar 'Fairway Ruby') (Accession numbers: X57294(embl),780766(gi)); CbVd.3 (Coleus blumei viroid 3-FR stem-loop RNA, from the Coleus blumei cultivar 'Fairway Ruby') (Accession numbers: X95290(embl), 1770105(gi))]

#### Hop latent viroid (HLVd)

30 [HLVd.1 (Accession numbers: X07397(embl), 60259(gi)); HLVd.2 ('thermomutant' T15) (Accession numbers: AJ290404(gb), 13872743(gi));

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HLVd.3 ('thermomutant' T40)(Accession AJ290405.1(gb), numbers: 13872744(gi)); HLVd.4 ('thermomutant' T50)(Accession numbers: AJ290406(qb), 13872745(gi)); HLVd.5 ('thermomutant' T59)(Accession numbers: AJ290406(gb), 13872746(gi)); HLVd.6 ('thermomutant' T61) (Accession numbers: AJ290408(gb) 13872747(gi)); HLVd.7 ('thermomutant' T75)(Accession numbers: AJ290409(gb), 13872748(gi)); HLVd.8 ('thermomutant' T92) (Accession numbers: AJ290410(gb), 13872749(gi)); HLVd.9 ('thermomutant' T218) (Accession numbers: AJ290411(gb), 13872750(gi)); HLVd.10 ('thermomutant' T229)(Accession numbers: AJ290412(gb), 13872751(gi))]

Australian grapevine viroid (AGVd) [AGVd.1 (Accession numbers: X17101(embl), 58574(gi))]

**Tomato planta macho viroid (TPMVd)** [TPMVd.1 (Accession numbers: K00817(gb))]

Coconut tinangaja viroid (CTiVd) [CTiVd.1 (Accession numbers: M20731(gb), 323414(gi))]

Tomato apical stunt viroid (TASVd) [TASVd.1 (Accession numbers K00818(gb), 335155(gi)); TASVd.2 (strain: indonesian) (Accession numbers: X06390(embl), 60650(gi)); TASVd.3(Tomato apical stunt viroid-S stem-loop RNA.) (Accession numbers: X95293(embl), 1771788(gi))]

Cadang-cadang coconut viroid (CCCVd) [CCCVd.1 (isolate baao 54, ccRNA 1 fast) (Accession numbers: J02049(gb), 323275(gi)); CCCVd.2 (isolate baao 54, ccRNA 1 fast) (Accession numbers: J02050(gb), 323276(gi)); CCCVd.3 (isolate baao 54, ccRNA 1 slow) (Accession numbers: J02051(gb), 323277(gi)); CCCVd.4 (isolates Ligao 14B, 620C, 191D and T1, ccRNA 1 fast) (Haseloff et al. *Nature* 299, 316-321 (1982)) CCCVd.5 (isolates Ligao T1, ccRNA 1 slow) (Haseloff et al. *Nature* 299, 316-321 (1982)); CCCVd.6 (isolates Ligao 14B, ccRNA 1 slow) (Haseloff et al. *Nature* 299, 316-321 (1982)); CCCVd.7 (isolate San Nasciso, ccRNA 1 slow) (Haseloff et al. *Nature* 299, 316-321 (1982))]

Citrus exocortis viroid (CEVd) [CEVd.1 (cev from gynura) (Accession numbers: J02053(gb), 323302(gi)); CEVd.2 (strain A) (Accession numbers:

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M34917(gb), 323305(gi)); CEVd.3 (strain de25)(Accession numbers: K00964(gb), 323303(gi)); CEVd.4 (strain de26) (Accession numbers: K00965(qb), 323304(qi)); CEVd.5 (CEV-JB) (Accession numbers: M30870(qb), 484119(gi)); CEVd.6 (CEV-JA) (Accession numbers: M30869(gb), 484118(gi)); CEVd.7 (Accession numbers: M30871(gb), 484117(gi)); CEVd.8 (CEV-A)(Accession numbers: M30868(gb), 484116(gi)); CEVd.9 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)) CEVd.10 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.11 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.12 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.13 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.14 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.15 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.16 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.17 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.18 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.19 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.20 (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.21 (cev-j classe B) (Visvader, J.E. and Symons, R.H. Nucleic Acids Res. 13, 2907-2920 (1985)); CEVd.22 (Grapevine viroid (GV)) (Accession numbers: Y00328(embl), 60645(gi)); CEVd.23 (CEVd-t) (Accession numbers: X53716(embl), 433503(gi)); CEVd.24 (CEVcls, isolate tomato hybrid callus) (Accession numbers: S67446(gb), 141247(gi)); CEVd.25 (CEV D-92) (Accession numbers: S67442(gb), 141248(gi)); CEVd.26 (CEVt, isolate tomato hybrid) (Accession numbers: S67441(gb), 141246(gi)); CEVd.27 (CEVt, isolate tomato)(Accession numbers: S67440(gb), 141245(gi)); CEVd.28 (CEVg, isolate Gynura) (Accession numbers: S67438(gb), 141244(gi)); CEVd.29 (CEVc, isolate citron)(Accession numbers: S67437(gb), 141243(gi)); CEVd.30 (strain CEVd-225) (Accession numbers: U21126(gb), 710360(qi)); CEVd.31 ( isolate broad bean, Vicia faba L.) (Accession numbers:

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S79831(gb),1181910(gi)); CEVd.32 (variant obtain after inoculation tomato with cevd.31) (Fagoaga et al. *J. Gen. Virol.* **76**, 2271-2277 (1995));

CEVd.33 (Fagoaga et al. *J. Gen. Virol.* **76**, 2271-2277 (1995)); CEVd.34 (Accession numbers: AF298177, 15419885(gi)); CEVd.35 (Accession numbers: AF298178, 15419886(gi)); CEVd.36 (Accession: AF428058) (Citrus exocortis viroid isolate 205-E-1 Uy, complete genome.); CEVd.37 (Accession: AF428059) (Citrus exocortis viroid isolate 205-E-2 Uy, complete genome.); CEVd.38 (Accession: AF428060) (Citrus exocortis viroid isolate 205-E-5 Uy, complete genome.); CEVd.39 (Accession: AF428061) (Citrus exocortis viroid isolate 205-E-7 Uy, complete genome.); CEVd.40 (Accession: AF428062) (Citrus exocortis viroid isolate 54-E-1 Uy, complete genome.); CEVd.41 (Accession: AF428063) (Citrus exocortis viroid isolate 54-E-3 Uy, complete genome.); CEVd.42 (Accession: AF428064) (Citrus exocortis viroid isolate 54-E-18 Uy, complete genome.); CEVd.43 (Accession: AF434678) (Citrus exocortis viroid, complete genome.)]

Columnea latent viroid (CLVd) [CLVd.1 (Accession numbers: X15663(embl), 58886(gi)); CLVd.2 (CLVd-N, individual isolate Nematanthus) (Accession numbers: M93686(gb), 323335(gi)); CLVd.3(Columnea latent viroid-B stemloop RNA) (Accession numbers: X95292(embl), 1770174(gi))]

Citrus bent leaf viroid (CBLVd) [CBLVd.1 (CVd-lb) (Accession numbers: M74065(gb), 323413(gi)); CBLVd.2 (strain CBLVd-225) (Accession numbers: U21125(gb), 710359(gi)); CBLVd.3 (viroid la genomic RNA, isolate: Jp) (Accession numbers: AB006734(dbj), 2815403(gi)); CBLVd.4 (viroid lb genomic RNA, isolate: P2) (Accession numbers: AB006735(dbj), 2815401(gi)); CBLVd.5 (viroid la genomic RNA) (Accession numbers: AB006736(dbj), 2815402(gi)); CBLVd.6 (Citrus Viroid la clone 17) (Accession numbers: AF040721(gb), 3273626(gi)); CBLVd.7 (Citrus Viroid la clone 18) (Accession numbers: AF040722(gb), 3273627(gi)); CBLVd.8 (Citrus bent leaf viroid isolate 201-1-1 Uy, complete genome.) (Accession: AF428052); CBLVd.9 (Citrus bent leaf viroid isolate 201-1-5 Uy, complete genome.)

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(Accession: AF428054); CBLVd.11 (Citrus bent leaf viroid isolate 205-1-1 Uy, complete genome.) (Accession: AF428055); CBLVd.12 (Citrus bent leaf viroid isolate 205-1-3 Uy, complete genome.) (Accession: AF428056); CBLVd.13 (Citrus bent leaf viroid isolate 205-1-4 Uy, complete genome.) (Accession: AF428057)]

Hop stunt viroid (HSVd) [HSVd.h1 (Japanese type strain) (Accession numbers: X00009(embl), 60684(gi)); HSVd.h2 (Japanese strain, variant 2) (Lee et al. Nucleic Acids Res. 16, 8708-8708 (1988)); HSVd.h3 (Korean strain) (Accession numbers: X12537(embl), 60421(gi)); HSVd.g1 (Grapevine viroid (GVVd), isolate SHV-g(GV)) (Accession numbers: M35717(gb), 325405(gi)); HSVd.g2 (strain: German cultivar Riesling) (Accession numbers: X06873(embl), 60422(gi)); HSVd.g3 (strain: isolated from Vitis vinifera Rootstock 5BB) (Accession numbers: X15330(embl), 60648(gi)); HSVd.g4 (isolate grapevine (HSVdg), variant la) (Accession numbers: X87924(embl), 897764(gi)); HSVd.g5 (isolate grapevine (HSVdg), variant lb) (Accession numbers: X87923(embl), 897765(gi)); HSVd.g6 (isolate grapevine (HSVdg), variant lc) (Accession numbers: X87925(embl), 897766(gi)); HSVd.g7 (isolate grapevine (HSVdq), variant ld)(Accession numbers: X87926(embl), 897767(gi)); HSVd.g8 (isolate grapevine (HSVdg), variant le) (Accession numbers: X87927(embl), 897768(gi)); HSVd.g9 (isolate grapevine (HSVdg), variant IIa) (Accession numbers:X87928(embl), 897769(gi)); HSVd.cit1 (variant 1, isolate HSV-cit) (Accession numbers: X06718(embl), 60646(gi)); HSVd.cit2 (variant 2, isolate HSV-cit) (Accession numbers: X06719(embl), 60647(gi)); HSVd.cit3 (HSV.citrus) (Accession numbers: X13838(embl), 60418(gi)); HSVd.cit4(Accession numbers: U02527(gb), 409021(qi)); HSVd.cit5 (Hsu et al. Virus Genes 9, 193-195 (1995)); HSVd.cit6 cit5 (Hsu et al. Virus Genes 9, 193-195 (1995)); HSVd.cit7 (isolate CVd-IIa or E819) (Accession numbers: AF131248(qb)); HSVd.cit8 (isolate CVd-IIb or Ca902) (Accession numbers: AF131249(gb)); HSVd.cit9 (isolate CVd-IIc or Ca905) (Accession numbers: AF131250(gb)); HSVd.cit10 (isolate Ca903) (Accession numbers: AF131251(gb)); HSVd.cit11 (isolate CA909) (Accession

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numbers: AF131252(gb)); HSVd.cit12 (cachexia isolate X-701-M) (Accession numbers: AF213483(gb), 12082502(gi)); HSVd.cit13 (cachexia isolate X-701-1) (Accession numbers: AF213484(gb), 12082503(gi)); HSVd.cit14 (cachexia isolate X-701-2) (Accession numbers: AF213485(gb), 12082504(gi)); HSVd.cit15 (cachexia isolate X-701-3) (Accession numbers: AF213486(gb), 12082505(gi)); HSVd.cit16 (cachexia isolate X-704-M) (Accession numbers: AF213487(gb), 12082506(gi)); HSVd.cit17 (cachexia isolate (Accession numbers: AF213488(gb), 12082507(gi)); HSVd.cit18 (cachexia isolate X-704-2) (Accession numbers: AF213489(gb), HSVd.cit19 (cachexia isolate X-704-3) (Accession numbers: AF213490(gb), 12082509(gi)); HSVd.cit20 (cachexia isolate X-707-M) (Accession numbers: AF213491(gb), 12082510(gi)); HSVd.cit21 (cachexia isolate (Accession numbers: AF213492(gb), 12082511(gi)); HSVd.cit22 (cachexia isolate X-707-2) (Accession numbers: AF213493(gb), 12082512(qi)); HSVd.cit23 (cachexia isolate X-707-3) (Accession numbers: AF213494(gb), 12082513(gi)); HSVd.cit24 (cachexia isolate X-707-4) (Accession numbers: AF213495(qb), 12082514(gi)); HSVd.cit25 (cachexia isolate X-712-M) (Accession numbers: AF213496(gb), 12082515(gi)); HSVd.cit26 (cachexia isolate X-712-1) (Accession numbers: AF213497(gb), 12082516(gi)); HSVd.cit27 (cachexia isolate X-712-2) (Accession numbers: AF213498(gb), 12082517(gi)); HSVd.cit28 (cachexia isolate X-712-3) (Accession numbers: AF213499(gb), 12082518(gi)); HSVd.cit29 (cachexia isolate X-715-M) (Accession numbers: AF213500(gb), 12082519(gi)); HSVd.cit30 (cachexia isolate X-715-1) (Accession numbers: AF213501(gb), HSVd.cit31 (cachexia isolate X-715-2) (Accession numbers: AF213502(gb), 12082521(gi)); HSVd.cit32 (CVd-lia (117)(Accession AF213503(gb), 12082522(gi)); HSVd.cit33 (isolate CVd-lla 17uy) (Accession numbers: AF359276(gb), 13991644(gi)); HSVd.cit34 (isolate CVd-lla 11uy) (Accession numbers: AF359275(gb), 13991643(gi)); HSVd.cit35 (isolate CVd-IIa 10uy) (Accession numbers: AF359274(gb), 13991642(gi)); HSVd.cit36 (isolate CVd-lb 10uy) (Accession numbers:

AF359273(gb), 13991641(gi)); HSVd.cit37 (isolate CVd-lb 5uy) (Accession numbers: AF359272(gb), 13991640(gi)); HSVd.cit38 (isolate CVd-lb 3uy) (Accession numbers: AF359271(gb), 13991639(gi)); HSVd.cit39 (isolate CVd-(Accession numbers: AF359270(gb), 13991638(qi)); 2uy) HSVd.cit40 (isolate CVd-IIa) (Accession numbers: X69519(embl), 2369773(gi)); 5 HSVd.cit41 (isolate CVd-IIb) (Accession numbers: X69518(embl),2369774(gi)); HSVd.cit42 (isolate CVd-lla 54-2-1) (Accession numbers: AF416554, 15811645(gi)); HSVd.cit43 (isolate CVd-IIa 54-2-2) (Accession numbers: AF416555, 15811646(gi)); HSVd.cit44 (isolate CVd-IIa 205-2-4) (Accession numbers: AF416556, 10 15811647(gi)); HSVd.cit45 (isolate CVd-lla 205-2-1) (Accession numbers: AF416557, 15811648(gi)); HSVd.p1 (HSV-peach (A9)) (Accession numbers: D13765(dbj), 221254(gi)); HSVd.p2 (HSV-plum and HSV-peach (AF) isolate) (Accession numbers: D13764(dbj), 221255(gi)); HSVd.p3 (cv. Jeronimo J-16 from Spain) (Accession numbers: Y09352(embl),1684696(gi)); HSVd.apr1 (cv. 15 Rouge de Roussillon from France) (Accession numbers: Y08438(embl), 2462494(gi)); HSVd.apr2 (unknown cultivar from Spain) (Accession numbers: Y08437 (embl), 2462495(gi)); HSVd.apr3 (cv. Bulida from Spain) (Accession numbers: Y09345(embl),1684690(gi)); HSVd.apr4 (cv. Bulida from Spain) (Accession numbers: Y09346(embl),1684691(gi)); HSVd.apr5 (cv. Bulida 20 d'Arques from Spain) (Accession numbers: Y09344(embl),1684692(gi)); (cv. HSVd.apr6 Pepito del Rubio from (Accession Spain) numbers:Y09347(embl), 1684697(gi)); HSVd.apr7 (cv. Pepito del Rubio from Spain) (Accession numbers: 09348(embl), 1684699(gi)); HSVd.apr8 (cv. Pepito del Rubio from Spain) (Accession numbers: Y09349(embl), 684698(gi)); 25 HSVd.apr9 (cv. Canino from Morocco) (Accession numbers: AJ297825(gb), 10944963(gi)); HSVd.apr10 (cv. Canino from Morocco) (Accession numbers: AJ297826(gb), 10944964(gi)); HSVd.apr11 (cv. Canino (Accession from Morocco) numbers: AJ297827(gb), 10944965(gi)); HSVd.apr12 (cv. Canino from Morocco) (Accession numbers: AJ297828(gb), 30 10944966(gi)); HSVd.apr13 (cv. Canino from Morocco) (Accession numbers:

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AJ297829(gb), 10944967(gi)); HSVd.apr14 (cv. Septik from Turkey) (Accession numbers: AJ297830(gb), 10944968(qi)); HSVd.apr15 (cv. Monaco bello from Cyprus) (Accession numbers: AJ297831(qb), 10944969(gi)); HSVd.apr16 (cv.Cafona from Cyprus) (Accession numbers: AJ297832(gb), 10944970(gi)); HSVd.apr17 (cv.Cafona from Cyprus) (Accession numbers: AJ297833(gb), 10944971(gi)); HSVd.apr18 (cv.Boccuccia spinosa from Cyprus) (Accession numbers: AJ297834(gb), 10944972(gi)); HSVd.apr19 (cv. Palumella from Cyprus) (Accession numbers: AJ297835(gb), 10944973(gi)); HSVd.apr20 (cv. Palumella from Cyprus) (ccession numbers: AJ297836(gb), 10944974(gi)); HSVd.apr21 (cv.Canino from Cyprus) (Accession numbers: AJ297837(qb), 10944975(gi)); HSVd.apr22 (cv.Kolioponlou from Greece) (Accession numbers: AJ297838(gb), 10944976(gi)); HSVd.apr23 (cv. Bebecou Paros from Greece) (Accession numbers: AJ297839(gb), 10944977(gi)); HSVd.apr24 (cv. Bebecou Paros from Greece) (Accession numbers: AJ297840(qb), 10944978(qi)); HSVd.c1 (Cucumber pale fruit viroid (CPFVd), isolate HSV-cucumber) (Accession numbers: X00524(embl), 60644(gi)); HSVd.c2 (Cucumber pale fruit viroid (CPFVd)) (Accession numbers: X07405(embl), 59015(gi)); HSVd.alm1 (Accession numbers: AJ011813(emb), 3738118(gi)); HSVd.alm2 (Accession numbers: AJ011814(emb), 3738119(gi)); HSVd. Citrus viroid II, complete genome (Accession number: AF434679)]. All these nucleotide sequences are herein incorporated by reference.

As will be immediately apparent from the above list, viroids are extremely prone to sequence variations, and such natural variants can also be used for the currently described methods and means, particularly if they retain the capacity to be transported to the nucleus, together with any operably linked RNA.

In addition to the natural variations in viroid nucleotide sequences, variants may be obtained by substitution, deletion or addition of particular nucleotides, and such variants may also be suitable for the currently described methods and WO 03/076619 PCT/AU03/00292

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means, particularly if they retain the capacity to be transported to the nucleus, together with any operably linked RNA.

Further, smaller RNA regions derived from the viroid nucleotide sequences, and variants thereof can be used for the current invention which are capable of being transported to the nucleus together with any operably linked RNA.

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The capacity of both smaller regions and variants derived from viroid nucleotide sequences to be transported to the nucleus of a host cell, such as a plant cell, can be determined using the assay described by Zhou et al. 2001, J. Gen Virology, 82, 1491-1497. Briefly, the assay comprises introducing a marker coding region, such as GFP, comprising an intervening sequence in the coding region of the marker gene, into the host cell by means of a viral RNA vector that replicates in the cytoplasm of the host cell. When a functional nuclear localization signal is introduced (conveniently inserted in the intervening sequence), the viral RNA vector comprising the marker gene is imported into the nucleus, where the intron can be removed and the spliced RNA returned to the cytoplasm. The spliced RNA can be detected by the translation into GFP protein, as well as by RNA analysis methods (e.g. RT-PCR) to confirm the absence of the intron from the spliced RNA molecules.

Furthermore, the human hepatitis delta RNA is a 1700 nt single stranded circular RNA which is very similar to the viroids of the PSTVd-type in that is localized in the nucleus, forming rod-like structures, and may also be used according to the invention.

In another embodiment of the invention, the largely double stranded RNA region comprises CUG, CAG, GAC OR GUC repeats. As used herein « trinucleotide repeats or CUG, CAG, GAC OR GUC repeats » are RNA molecules comprising a number of CUG, CAG, GAC OR GUC trinucleotides. Preferably, the CUG trinucleotides are repeated without intervening sequences,

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although short regions of 1 to 20-30 nucleotides not consisting of CUG trinucleotides may be present occasionally between the CUG trinucleotide repeats. Preferably, the CUG repeats comprise a number of CUG trinucleotide exceeding 35 copies or 44 copies such as any number between 50 and 2000 copies. Conveniently the copy number of the CUG triplets should not exceed 100 or 150. It is expected that CAG, GAC or GUC repeats may be used to similar effect.

Without intending to limit the invention to a particular mode of action, it is taught that such trinucleotide repeats repeats form rod-like structures by imperfect base-pairing which function as nuclear retention signal, possibly by sterically blocking RNA export through nuclear pores, as well as activate double stranded RNA dependent protein kinase PKR [ Davis et al , 1997 Proc. Natl. Acad. Sci. 94, 7388-7393; Tian et al. 2000 RNA 6, 79-87; Koch and Lefert 1998 J. Theor. Biol. 192, 505-514).

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CUG repeats may be particularly suited to increase the efficiency of antisensemediated gene silencing when the RNA molecules comprising such CUG repeats can be delivered to the nucleus of the host cell e.g. through transcription of a chimeric gene encoding such RNA, as hereinafter described.

Although the largely double stranded RNA region such as the PSTVd-type viroid derived nuclear location signals or the trinucleotide repeats can conveniently be located at the 3' end of the target specific antisense RNA, it is expected that the location of the largely double stranded RNA is of little importance. Hence, largely double stranded RNA regions may also be located at the 5' end of the RNA molecule preferably at the 3' end or even in the middle of such an RNA molecule.

30 It was also unexpectedly found that the efficiency of antisense-mediated downregulation of gene expression, wherein the antisense RNA was operably

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linked to a largely double stranded RNA region, could be further enhanced by inclusion of an intron sequence in the RNA molecule provided to the host cell. Again, the location of the intron in the RNA molecule with respect to both the target specific nucleotide sequence as well as the the largely double-stranded RNA region is expected to have little effect on the efficiency. In fact, it is expected that the largely double stranded RNA region may be located within the intron sequence.

As used herein, an "intron" or intervening sequence is used to refer to a DNA region within a larger transcribed DNA region, which is transcribed in the nucleus to yield an RNA region which is part of a larger RNA, however, said RNA region corresponding to intro sequence is removed from the larger RNA when transferred to the cytoplasm. The corresponding RNA is also referred to as an intron or intervening sequence. Intron sequences are flanked by splicing sites, and synthetic introns may be made by joining appropriate splice sites to basically any sequence, having an approriate branching point. Introns or intervening sequences which are located in 5'UTR, coding region or 3'UTR may be used.

Intervening sequences or introns should preferably be capable of being spliced in the eukaryotic host cells, although the presence of intervening sequences which can no longer be spliced, e.g. because their guide sequences have been altered or mutated, may even further increase the efficiency of the chimeric RNA molecules to down regulate the expression of a target gene. In one embodiment of the invention, the intron is essentially identical in sequence to the *Flaveria trinervia* pyruvate orthophosphate dikinase 2 intron 2 (pdk2 intron) and may comprise the sequence of SEQ ID No 9. Other examples of plant introns include the catalase intron from Castor bean (Accession number AF274974), the Delta12 desaturase (Fad2) intron from cotton (Accession number AF331163), the Delta 12 desaturase (Fad2) intron from *Arabidopsis* (Accession number AC069473), the Ubiquitin intron from maize (Accession

number S94464), the actin intron from rice. Other examples of mamalian virus introns include the intron from SV40. Examples of fungal introns include the intron from the triose phosphate isomerase gene from Aspergillus.

It was also unexpectedly found that further introduction of a sense RNA molecule with a target-gene specific region corresponding to the target gene specific region of the antisense RNA molecule already present in the cell of the eukaryotic organism, further increased the efficiency of the downregulation of the expression of the target gene. The same efficiency of downregulation of the expression of a target gene could be observed if the sense RNA molecule was provided with a largely double stranded RNA region as herein described. Sense RNA molecule was provided to a cell of a eukaryotic host organism simultaneously with an antisense RNA molecule capable of forming a double stranded region by basepairing with the sense RNA molecule.

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Thus, in another embodiment of the invention a method is provided for down regulating the expression of a target gene in cells of a eukaryotic organisms, comprising the steps of

- a) providing the cells of the eukaryotic organism with a first and second chimeric RNA molecule, wherein
  - i) the first chimeric RNA molecule comprises an antisense target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene;

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ii) the second chimeric RNA molecule comprises a sense target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity to the complement of the first chimeric RNA molecule;

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iii) the first and second chimeric RNA are capable of basepairing at least between the 19 consecutive nucleotides of the first chimeric

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RNA and the 19 consecutive nucleotides of the second chimeric RNA; and

- iv) wherein either the first or the second chimeric RNA molecule comprises a largely double stranded RNA region operably linked to the antisense target-specific RNA region or to the sense target-specific RNA region; and
- b) identifying those eukaryotic organisms wherein the expression of the target gene is down regulated.

In another specific embodiment, both the first and second chimeric RNA molecule comprise a largely double stranded region. Specific embodiments for the largely double stranded RNA region and target gene-specific antisense RNA are as described elsewhere in this application. Specific embodiments for the sense RNA region are similar to the specific embodiments for the antisense RNA region.

Conveniently, the antisense or sense RNA molecules comprising a largely double stranded RNA region as herein described may be provided to the eukaryotic host cell or organism by introduction and possible integration of a chimeric gene, transcription of which yields such an antisense or sense RNA. Thus the invention is also aimed at providing such a chimeric gene comprising

- a promoter or a promoter region which is capable of being expressed in cells of the eukaryotic organism of interest; operably linked to a DNA region which when transcribed yields an antisense RNA molecule comprising
  - a target-gene specific antisense nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene; or
  - a target-gene specific sense nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene

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identity with 19 consecutive nucleotides from the nucleotide sequence of the target gene; operably linked to

- a largely double stranded RNA region as herein described; and optionally
- a transcription termination and polyadenylation region suitable for the eukaryotic cell of choice.

As used herein, the term "promoter" denotes any DNA which is recognized and bound (directly or indirectly) by a DNA-dependent RNA-polymerase during initiation of transcription. A promoter includes the transcription initiation site, and binding sites for transcription initiation factors and RNA polymerase, and can comprise various other sites (e.g., enhancers), at which gene expression regulatory proteins may bind.

The term "regulatory region", as used herein, means any DNA, that is involved in driving transcription and controlling (i.e., regulating) the timing and level of transcription of a given DNA sequence, such as a DNA coding for a protein or polypeptide. For example, a 5' regulatory region (or "promoter region") is a DNA sequence located upstream (i.e., 5') of a coding sequence and which comprises the promoter and the 5'-untranslated leader sequence. A 3' regulatory region is a DNA sequence located downstream (i.e., 3') of the coding sequence and which comprises suitable transcription termination (and/or regulation) signals, including one or more polyadenylation signals.

In one embodiment of the invention the promoter is a constitutive promoter. In another embodiment of the invention, the promoter activity is enhanced by external or internal stimuli (inducible promoter), such as but not limited to hormones, chemical compounds, mechanical impulses, abiotic or biotic stress conditions. The activity of the promoter may also regulated in a temporal or spatial manner (tissue-specific promoters; developmentally regulated promoters).

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In a particular embodiment of the invention, the promoter is a plant-expressible promoter. As used herein, the term "plant-expressible promoter" means a DNA sequence which is capable of controlling (initiating) transcription in a plant cell. This includes any promoter of plant origin, but also any promoter of non-plant origin which is capable of directing transcription in a plant cell, i.e., certain promoters of viral or bacterial origin such as the CaMV35S (Hapster et al., 1988), the subterranean clover virus promoter No 4 or No 7 (WO9606932), or T-DNA gene promoters but also tissue-specific or organ-specific promoters including but not limited to seed-specific promoters (e.g., WO89/03887), organprimordia specific promoters (An et al., 1996), stem-specific promoters (Keller et al., 1988), leaf specific promoters (Hudspeth et al., 1989), mesophyl-specific promoters (such as the light-inducible Rubisco promoters), root-specific promoters (Keller et al., 1989), tuber-specific promoters (Keil et al., 1989), vascular tissue specific promoters (Peleman et al., 1989), stamen-selective promoters (WO 89/10396, WO 92/13956), dehiscence zone specific promoters ( WO 97/13865) and the like.

In another particular embodiment of the invention, the promoter is a fungus-expressible promoter. As used herein, the term "fungus-expressible promoter" means a DNA sequence which is capable of controlling (initiating) transcription in a fungal cell such as but not limited to the *A. nidulans trp*C gene promoter, or the inducible *S. cerevisiae* GAL4 promoter.

In yet another particular embodiment of the invention, the promoter is a animal-expressible promoter. As used herein, the term "animal-expressible promoter" means a DNA sequence which is capable of controlling (initiating) transcription in an animal cell and including but not limited to SV40 late and early promoters, cytomegalovirus CMV-IE promoters, RSV-LTR promoter, SCSV promoter, SCBV promoter and the like.

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The antisense or sense RNA molecules useful for the invention may also be produced by in vitro transcription. To this end, the promoter of the chimeric genes according to the invention may be a promoter recognized by a bacteriophage single subunit RNA polymerase, such as the promoters recognized by bacteriophage single subunit RNA polymerase such as the RNA polymerases derived from the E. coli phages T7, T3,  $\phi$ I,  $\phi$ II, W31, H, Y, A1, 122, cro, C21, C22, and C2; Pseudomonas putida phage gh-1; Salmonella typhimurium phage SP6; Serratia marcescens phage IV; Citrobacter phage VillI; and Klebsiella phage No.11 [Hausmann, Current Topics in Microbiology and Immunology, 75: 77-109 (1976); Korsten et al., J. Gen Virol. 43: 57-73 (1975); Dunn et al., Nature New Biology, 230: 94-96 (1971); Towle et al., J. Biol. Chem. 250: 1723-1733 (1975); Butler and Chamberlin, J. Biol. Chem., 257: 5772-5778 (1982)]. Examples of such promoters are a T3 RNA polymerase specific promoter and a T7 RNA polymerase specific promoter, respectively. A T3 promoter to be used as a first promoter in the CIG can be any promoter of the T3 genes as described by McGraw et al, Nucl. Acid Res. 13: 6753-6766 (1985). Alternatively, a T3 promoter may be a T7 promoter which is modified at nucleotide positions -10, -11 and -12 in order to be recognized by T3 RNA polymerase [(Klement et al., J. Mol. Biol. 215, 21-29(1990)]. A preferred T3 promoter is the promoter having the "consensus" sequence for a T3 promoter, as described in US Patent 5,037,745. A T7 promoter which may be used according to the invention, in combination with T7 RNA polymerase, comprises a promoter of one of the T7 genes as described by Dunn and Studier, J. Mol. Biol. 166: 477-535 (1983). A preferred T7 promoter is the promoter having the "consensus" sequence for a T7 promoter, as described by Dunn and Studier (supra).

The antisense or sense RNA can be produced in large amounts by contacting the acceptor vector DNA with the appropriate bacteriophage single subunit RNA polymerase under conditions well known to the skilled artisan. The so-produced antisense or sense RNA can then be used for delivery into cells

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prone to gene silencing, such as plant cells, fungal cells or animal cells. Antisense RNA may be introduced in animal cells via liposomes or other transfection agents (e.g. Clonfection transfection reagent or the CalPhos Mammalian transfection kit from ClonTech) and could be used for methods of treatment of animals, including humans, by silencing the appropriate target genes. Antisense or sense RNA can be introduced into the cell in a number of different ways. For example, the antisense or sense RNA may be administered by microinjection, bombardment by particles covered by the antisense or sense RNA, soaking the cell or organisms in a solution of the antisense or sense RNA, electroporation of cell membranes in the presence of antisense or sense RNA, liposome mediated delivery of antisense or sense RNA and transfection mediated by chemicals such as calcium phosphate, viral infection, transformation and the like. The antisense or sense RNA may be introduced along with components that enhance RNA uptake by the cell, stabilize the annealed strands, or otherwise increase inhibition of the target gene. In the case of a whole animal, the antisense or sense RNA is conveniently introduced by injection or perfusion into a cavity or interstitial space of an organism, or systemically via oral, topical, parenteral (including subcutaneous, intramuscular or intravenous administration), vaginal, rectal, intranasal, ophthalmic, or intraperitoneal administration. The antisense or sense RNA may also be administered via an implantable extended release device.

The chimeric genes according to the invention capable of producing antisense or sense RNA may also be equipped with any prokaryotic promoter suitable for expression of the antisense or sense RNA in a particular prokaryotic host. The prokaryotic host can be used as a source of antisense and/or sense RNA, e.g. by feeding it to an animal, such as a nematode or an insect, in which the silencing of the target gene is envisioned and monitored by reduction of the expression of the reporter gene. In this case, it will be clear that the target gene and reporter genes should be genes present in the cells of the target eukaryotic organism and not of the prokaryotic host organism. The antisense and sense

RNA according to the invention or chimeric genes capable of yielding such antisense or sense RNA molecules, can thus be produced in one host organism, be administered to a another target organisms (e.g. through feeding, orally administring, as a naked DNA or RNA molecule or encapsulated in a liposome, in a virus particle or attentuated virus particle, or on an inert particle etc.) and effect reduction of gene expression in the target gene or genes in another organism.

Suitable transcription termination and polyadenylation region include but are not limited to the SV40 polyadenylation signal, the HSV TK polyadenylation signal, the nopaline synthase gene terminator of *Agrobacterium tumefaciens*, the terminator of the CaMV 35S transcript, terminators of the subterranean stunt clover virus, the terminator of the *Aspergillus nidulans* trpC gene and the like.

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The invention also aims at providing the antisense and sense RNA molecules, which may be obtained by transcription from these chimeric genes, and which are useful for the methods according to the invention.

It is another object of the invention to provide eukaryotic cells, and eukaryotic non-human organisms containing the antisense RNA molecules of the invention, or containing the chimeric genes capable of producing the antisense RNA molecules of the invention. In a preferred embodiment the chimeric genes are stably integrated in the genome of the cells of the eukaryotic organism.

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It is also an object of the invention to provide eukaryotic cells and eukaryotic non-human organisms containing simultaneously sense and antisense RNA molecules of which one or both of the RNA molecules comprise a largely double stranded RNA region, or chimeric genes encoding such RNA molecules.

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In another embodiment, the chimeric genes of the invention may be provided on a DNA molecule capable of autonomously replicating in the cells of the eukaryotic organism, such as e.g. viral vectors. The chimeric gene or the antisense or sense RNA may be also be provided transiently to the cells of the eukaryotic organism.

Introduction of chimeric genes (or RNA molecules) into the host cell can be accomplished by a variety of methods including calcium phosphate transfection, DEAE-dextran mediated transfection, electroporation, microprojectile bombardment, microinjection into nuclei and the like.

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Methods for the introduction of chimeric genes into plants are well known in the art and include *Agrobacterium*-mediated transformation, particle gun delivery, microinjection, electroporation of intact cells, polyethyleneglycol-mediated protoplast transformation, electroporation of protoplasts, liposome-mediated transformation, silicon-whiskers mediated transformation etc. The transformed cells obtained in this way may then be regenerated into mature fertile plants.

Transgenic animals can be produced by the injection of the chimeric genes into the pronucleus of a fertilized oocyte, by transplantation of cells, preferably uindifferentiated cells into a developing embryo to produce a chimeric embryo, transplantation of a nucleus from a recombinant cell into an enucleated embryo or activated oocyte and the like. Methods for the production of trangenic animals are well established in the art and include US patent 4, 873, 191; Rudolph et al. 1999 (Trends Biotechnology 17:367-374); Dalrymple et al. (1998) Biotechnol. Genet. Eng. Rev. 15: 33-49; Colman (1998) Bioch. Soc. Symp. 63: 141-147; Wilmut et al. (1997) Nature 385: 810-813, Wilmute et al. (1998) Reprod. Fertil. Dev. 10: 639-643; Perry et al. (1993) Transgenic Res. 2: 125-133; Hogan et al. Manipulating the Mouse Embryo, 2<sup>nd</sup> ed. Cold Spring Harbor Laboratory press, 1994 and references cited therein.

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Gametes, seeds, embryos, progeny, hybrids of plants or animals comprising the chimeric genes of the present invention, which are produced by traditional breeding methods are also included within the scope of the present invention.

The methods and means described herein, can be applied to any eukaryotic organism in which gene-silencing takes place, including but not limited to plants (such as corn, wheat, potato, sunflower, turf grasses, barley, rye, tomato, sugar cane, safflower, cotton, *Arabidopsis*, rice, Brassica plants, vegetables, soybeans, tobacco, trees, flax, palm trees, peanuts, beans, etc.) invertebrate animals (such as insects, shellfish, molluscs, crustaceans such as crabs, lobsters and prawns) vertebrate animals (fish, avian animals, mammals, humans), yeast and fungi amongst others.

The following non-limiting Examples describe method and means for enhanced antisense RNA mediated silencing of the expression of a target gene in eukaryotic cell or combined sense/antisense RNA mediated target gene silencing.

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Unless stated otherwise in the Examples, all recombinant DNA techniques are carried out according to standard protocols as described in Sambrook *et al.* (1989) *Molecular Cloning: A Laboratory Manual*, Second Edition, Cold Spring Harbor Laboratory Press, NY and in Volumes 1 and 2 of Ausubel *et al.* (1994) *Current Protocols in Molecular Biology, Current Protocols*, USA. Standard materials and methods for plant molecular work are described in *Plant Molecular Biology Labfax* (1993) by R.D.D. Croy, jointly published by BIOS Scientific Publications Ltd (UK) and Blackwell Scientific Publications, UK. Other references for standard molecular biology techniques include Sambrook and Russell (2001) *Molecular Cloning: A Laboratory Manual*, Third Edition, Cold Spring Harbor Laboratory Press, NY, Volumes I and II of Brown (1998) *Molecular Biology LabFax*, Second Edition, Academic Press (UK). Standard materials and methods for polymerase chain reactions can be found in

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Dieffenbach and Dveksler (1995) *PCR Primer: A Laboratory Manual*, Cold Spring Harbor Laboratory Press, and in McPherson at al. (2000) *PCR - Basics: From Background to Bench*, First Edition, Springer Verlag, Germany.

- Throughout the description and Examples, reference is made to the following sequences:
  - SEQ ID N°1: oligonucleotide primer for the amplication of the RG1 PSTVd
  - SEQ ID N°2: oligonucleotide primer for the amplication of the RG1 PSTVd
- SEQ ID N°3: nucleotide sequence of the genome of PSTVd RG1
  - SEQ ID N°4: nucleotide sequence of genome of the Australian grapevine viroid
  - SEQ ID N°5: nucleotide sequence of the genome of the Coconut tinangaja viroid
  - SEQ ID N° 6: nucleotide sequence of the genome of the Tomato planta macho viroid
  - SEQ ID N°7: nucleotide sequence of the genome of the Hop latent viroid
  - SEQ ID N°8: nucleotide sequence of the genome of the Tomato apical stunt viroid
  - SEQ ID N°9: nucleotide sequence of the pdk2 intron
- 20 SEQ ID N°10: nucleotide sequence of the EIN2 cDNA
  - SEQ ID N° 11: nucleotide sequence the genomic EIN2 clone
  - SEQ ID N° 12: oligonucleotide primer 1 for the amplication of the EIN2 part used in the constructs in the Examples
  - SEQ ID N° 13: oligonucleotide primer 2 for the amplification of the EIN2 part used in the constructs in the Examples.
  - SEQ ID N° 14: pTSVd sequence in pMBW491.
  - SEQ ID N° 15: pTSVd sequence in pMBW489 (with 10 nt deletion).

## **Examples**

Example 1: Construction of the different plant lines containing different chimeric genes used.

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As an example target gene to down-regulate the expression using the various constructs, the EIN2 gene from *Arabidopsis thaliana* was chosen. The down-regulation of the expression of the EIN2 gene can easily be visualized by germinating seeds on MS-ACC medium (containing aminocyclopropane-1-carboxylic acid (ACC)) and incubating either in the dark or in light.

Dark-grown EIN2 silenced seedlings grown in the dark have a longer hypocotyl and a more developed root system compared to wt seedlings, whereas EIN2 silenced seedlings grown in light can be differentiated from the wt seedlings by their larger cotyledon size (see Figure 3)

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The EIN2 nucleotide sequence to be used in the different constructs in sense or antisense orientation was amplified by PCR using oligonucleotide primers with a nucleotide sequence as represented in SEQ ID N° 12 and 13 using genomic DNA (nucleotide sequence see SEQ ID N° 11) or cDNA (nucleotide sequence see SEQ ID N° 10) as template DNA. The amplication of the genomic EIN2 sequence part (gEIN2) resulted in a PCR fragment with the nucleotide sequence of SEQ ID N° 11 from the nucleotide at position 538 to the nucleotide at position 1123 and contains two native introns of the EIN2 gene.

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The gEIN2 fragment was cloned as a Kpnl/Clal fragment into pART7 (Gleave, 1992 Plant. Mol. Biol. 20: 1203-1207), resulting in pMBW313 and the 35S promoter-gEIN2<sub>sense</sub>-OCS3' cassette was cloned into pART27 (Gleave 1992 supra) at the Notl site to result in pMBW353.

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A similar fragment (cEIN2) was amplified by PCR using EIN2 cDNA (SEQ ID N° 10) as template and the same pair of primers as for gEIN2. The cEIN2 fragment was digested with BamHI/Clal and cloned into pSHUTTLE (Wang et

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al., 1998 Acta Hort. 461: 401-407) at the same sites, giving pMBW310. The cEIN2 fragment was then excised from pMBW310 with Xbal and cloned into the Xbal site of pART7, forming pMBW351. From this intermediate vector the 35S-EIN2antisense-OCS3' cassette was excised and cloned into pWBVec2A (Wang et al. 1998, supra) at the Notl site, resulting in pMBW360.

A full length sequence of the PSTVd strain RG1 (SEQ ID N° 3) was amplified from a cDNA using oligonucleotides with the nucleotide sequence of SEQ ID N°1 and SEQ ID N°2. The resulting PCR fragment was digested with BgllI and cloned into the BamHI site of pMBW313, resulting in pMBW345, from which the 35S-gEIN2-PSTVd-OCS3' cassette was excised and cloned into pART27 at the NotI site resulting in pMBW355.

For pMBW359 the PCR amplified PSTVd sequence was digested with BgIII and cloned into the BamHI site of pMBW310, giving pMBW346, from which the cEIN2antisense-PSTVd sequence was excised with XbaI and cloned into the XbaI site of pHANNIBAL (Wesley et al. 2001), forming pMBW349. The 35S-pdk2-cEIN2antisense-PSTVd-OCS3' cassette was then cloned into pWBVec2a at the NotI site forming pMBW359. The cEIN2antisense PSTVd fragment was also cloned into pWBVec2a to yield pMBW357.

The EIN2 cDNA fragment was excised form pMBW310 with EcoRV/BamHI, blunted by Pfu treatment and ligated into the BamHI site (also Pfu treated) of pKANNIBAL (Wesley et al. 2001). Plasmids having the cEIN2 in both orientations with respect to the 35S promoter were recovered and named pMWB401 (antisense) and pMBW404 (sense orientation).

For pLMW37, pLMW38, pLMW39, and pLMW40 the cEIN2 fragment was inserted in sense or antisense orientation upstream or downstream of an inverted repeat of the PSTVd sequence. To this end, a partial PSTVd sequence (SEQ ID N° 3 from the nucleotide at position 16 to the nucleotide at position

355) was cloned upstream of the pdk intron in inverse orientation with regard to the complete copy of the PSTVd genome.

The different constructs are schematically represented in Figure 2.

Example 2 : <u>Analysis of expression of the EIN2 gene in transgenic *Arabidopsis* lines comprising the different chimeric genes of Example 1.</u>

The chimeric constructs represented in Figure 2 were introduced into *Agrobacterium* tumefaciens using conventional methods and the resulting *Agrobacterium* strains were used to introduce the chimeric genes into *Arabidopsis* ecotype Landsberg erecta through the dipping method. Transgenic lines were selected on 15 mg/L hygromycin or 50 mg/L kanamycin as the selective agent. T1 opr F1 seed was collected and assayed for EIN2 silencing.

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To this end, the seed was plated on MS medium containing  $50\mu$ M ACC. The plates were sealed tightly with parafilm and kept either under light or in the dark. Silencing was scored by looking at the size of roots and cotyledons (incubation in the light) or by looking at the size of roots or hypocotyls (incubation in the dark). In EIN2 silenced lines, the roots or hypocotyls are significantly longer, and the cotyledons are significantly larger than in wt lines grown under the same conditions.

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Seed from primary transformants was plated on MS-ACC medium, sealed with Parafilm, kept at 4C for 0-2 overnights, and then moved to growth room and kept either under light or in the dark. Silencing of the EIN2 gene was scored by examining the size of the roots and cotyledons (for those germinating under light) or the size of hypocotyls (for those in the dark). Significant or strongsilencing means long roots or hypocotyls, while weak silencing means bigger cotyledons but short roots or hypocotyls. The results are summarized in Table 1.

Table 1: Summary of the efficiency of EIN2 silencing in A. thaliana plants transformed with various EIN2 constructs.

Construct	Sr	nort description	transgenic lines	# strong silencing	# weak silencing	Frequency of silencing
PMBW360	0	EIN2 antisense	23	2	5	30%
PMBW401	0	EIN2 antisense	20	0	3	15%
	0	Pdk intron				
PMBW357	0	EIN2 antisense	17	3	5	47%
	0	PSTVd				
PMBW359	0	EIN2 antisense	22	10	6	73%
	0	PSTVd				
	0	Pdk intron	·			
PMBW353	0	EIN2 sense	19	2	3	26%
	0	Native introns				
PMBW355	0	EIN2 sense	17	1	1	12%
	0	Native introns				
	0	PSTVd				
PMBW404	0	EIN 2 sense	20	3	2	25%
	0	PDK intron				
PLMW37	0	EIN2 sense	19	0	0	0
	0	Pdk intron				
	0	PSTVd repeat				
PLMW38	0	EIN2 antisense	10	1	2	30
	0	Pdk intron				
	0	PSTVd repeat				
PLMW39	٥	EIN2 sense	17	0	0	0
	0	Pdk intron				
	0	PSTVd repeat				
PLMW40	0	EIN2 antisense	20	2	5	35%
	0	Pdk intron				
	0	PSTVd repeat				

Example 3: Analysis of expression of the EIN2 gene in Arabidopsis lines obtained by crossing of the transgenic Arabidopsis lines comprising the different chimeric genes of Example 1.

By cross-pollination between the *Arabidopsis* lines MBW353, MBW355, MBW359, MBW360 new lines were obtained containing simultaneously sense and antisense EIN2 constructs. These new lines were analyzed in a similar way

as described in Example 2. The results are summarized in Table 2. Plants wherein at least one of the transgenes contained a PSTVd sequence were very efficiently silenced.

Table 2. Summary of the efficiency of EIN2 silencing in A. thaliana plants comprising different combination of sense and antisense EIN2 constructs.

Line	Short description	N° of lines tested	N° of lines silenced	Frequency of silencing
MBW353 X MBW360	<ul><li>EIN2 sense</li><li>Native introns</li><li>And</li><li>EIN2 antisense</li></ul>	7	2	28.5%
MBW353 X MBW359	<ul> <li>EIN2 sense</li> <li>Native introns</li> <li>And</li> <li>EIN2 antisense</li> <li>PSTVd</li> <li>Pdk intron</li> </ul>	3	3	100%
MBW355 X MBW360	<ul><li>EIN2 sense</li><li>Native introns</li><li>PSTVd</li><li>And</li><li>EIN2 antisense</li></ul>	5	4	80%
MBW355 X MBW359	<ul> <li>EIN2 sense</li> <li>Native introns</li> <li>PSTVd</li> <li>And</li> <li>EIN2 antisense</li> <li>PSTVd</li> <li>Pdk intron</li> </ul>	11	9	81.8%

- Example 4: Construction of different chimeric genes for mediating gene silencing of a GFP gene in mammalian cells and analysis in CHO cells.
- As an example target gene to down-regulate the expression in mammalian cells, the humanized GFP coding region, expressed under control of a CMV promoter region, and followed by a SV40 polyadenylation signal was chosen (pCI-GFP)
- Different experimetal silencing constructs were constructed, having either the GFP coding region cloned in sense (as in pMBW493, pMBW494 and pMBW497) or antisense orientation (as in pMBW489, pMBW491 or pMBW496) with regard to the CMV promoter region.
- Plasmids pMBW493 and pMBW489 contained downstream of the GFP coding region, but upstream of the SV40 polyadenylation signal, a nucleotide sequence corresponding to a PSTVd sequence but with a 10 nt deletion (SEQ ID No 15). This deletion has an impact on the predicted secondary structure (see Fig 5).

Plasmids pMBW494 and pMBW491 contained downstream of the GFP coding region, but upstream of the SV40 polyadenylation signal, a nucleotide sequence corresponding to a PSTVd sequence of SEQ ID No 14 without the 10 nt deletion.

- Plasmids pMBW497 and pMBW496 contained downstream of the GFP coding region, but upstream of the SV40 polyadenylation signal, a nucleotide sequence comprising 60 CUG trinucleotide repeats.
- The different experimental plasmids were introduced (at different concentrations) into CHO cells in combination with a plasmid comprising the

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GFP expressing chimeric gene (Table 3; entries 1 to 18). Since the GFP construct is a functional sequence in the sense constructs, sense GFP containing experimental constructs were also introduced without the extra GFP expressing chimeric gene; to estimate the GFP expression by these constructs alone (Table 3; entries 19 to 30). Further, combinations of antisense and sense experimental constructs were introduced in CHO cells, at different concentrations(Table 3; entries 31 to 42). As a control, the chimeric GFP expression construct (pCi-GFP) was introduced alone into CHO cells.

After 24 hrs or 48 hrs, the cells were assayed for GFP expression. Average counts and standard deviations are represented in Table 3.

The antisense GFP constructs pMBW491, pMBW496 and pMBW489 that carry the pTSVd or CUG repeat sequences resulted in a significant reduction of the expression of the GFP gene.

Interestingly, pMWB489 in which the PSTVd sequence contains a 10 nt deletion, resulted in slower and lower degrees of GFP silencing than pMWB491, which contains an intact PSTVd sequence.

:	Experimental DNA	Target DNA	Remarks on Experimental DNA	Average	Standard	Average	Standard
· · · · -				count (24 hr)	deviation	count (48 hr)	deviation
-	0.1µg pMBW89	0.3µg GFP	Antisense+PSTVd (deletion)	3626	506	9028	1468
2	0.3µg pMBW 89	0.3µg GFP		3521	41	6468	3522
က	0.7µg pMBW 89	0.3µg GFP		3167	1348	1096	2191
4	0.1µg pMBW 91	0.3µg GFP	Antisense+PSTVd	3585	98	5908	623
ည	0.3µg pMBW 91	0.3µg GFP		748	128	1426	332.3
9	0.7µg pMBW 91	0.3µg GFP		23	25	1637	70
7	0.1µg pMBW 96	0.3µg GFP	Antisense + CUG repeats	3217	467	5221	4700
æ	0.3µg pMBW 96	0.3µg GFP	,	2907	107	3272	0
6	0.7µg pMBW 96	0.3µg GFP		181	92	1433	466
10	0.1µg pMBW 93	0.3µg GFP	Sense + PSTVd (deletion)	5815	313	16482	470
Ξ	0.3µg pMBW 93	0.3µg GFP		10453	1555	15810	1067
12	0.7µg pMBW 93	0.3µg GFP		12718	5423	10666	949
13	0.1µg pMBW 94	0.3µg GFP	Sense+PSTVd	9166	1269	15023	263
14	0.3µg pMBW 94	0.3µg GFP		12719	3894	6699	94
15	0.7µg pMBW 94	0.3µg GFP		1009	658	13133	824
16	$0.1\mu g pMBW 97$	0.3µg GFP	Sense+CUG repeats	6414	1367	15795	178
17	0.3µg pMBW 97	0.3µg GFP		3596	20	10235	770
18	0.7µg pMBW 97	0.3µg GFP		729	295	13171	2868
19	0.1µg pMBW 93	None	Sense + PSTVd (deletion)	1216	15	3692	142
50	0.3µg pMBW 93	None		6022	1293	9341	273
21	0.5µg pMBW 93	None		6795	3235	11466	2541
22	0.7µg pMBW 93	None		12002	763	10316	1523
23	0.1µg pMBW 94	None	Sense+PSTVd	2121	594	5417	777
24	0.3µg pMBW 94	None		5671	5096	9317	743
25	0.5µg pMBW 94	None		6349	3253	7842	337
56	0.7µg pMBW 94	None		1785	729	15574	2208
27	0.1µg pMBW 97	None	Sense+CUG repeats	4448	626	6064	289
28	0.3µg pMBW 97	None		487	83	7922	194
83	0.5µg pMBW 97	None		522	223	7481	999
30	0.7μg pMBW 97	None		270	159	8980	1154

	Experimental DNA	Target DNA	Remarks on Experimental DNA	Average	Standard	Average	Standard
				count	deviation	count	deviation
				(24 hr)		(48 hr)	
31	$0.1\mu g pMBW 93 + 0.1\mu g pMBW 91$	None	Sense + PSTVd (deletion) and	1189	148	2331	815
32	0.3µg pMBW 93 + 0.3µg pMBW 91	None	Antisense+PSTVd	695	83	3101	533
33	$0.5\mu g pMBW 93 + 0.5\mu g pMBW 91$	None		111	117	3758	1583
34	$0.3\mu g \text{ pMBW } 93 + 0.1\mu g \text{ pMBW } 91$	None		1811	1304	5301	23
32	$0.3\mu g pMBW 93 + 0.3\mu g pMBW 91$	None		312	171	4972	401
36	$0.3\mu g pMBW 93 + 0.7\mu g pMBW 91$	None		14.	20	2896	1075
37	$0.1\mu g pMBW 97 + 0.1\mu g pMBW 96$	None	Antisense+CUG repeats and	3841	929	2945	341
38	$0.3\mu g pMBW 97 + 0.3\mu g pMBW 96$	None	Sense+CUG repeats	1018	401	3236	822
39	$0.5\mu g pMBW 97 + 0.5\mu g pMBW 96$	None		1262	241	6730	583
40	$0.3\mu g pMBW 97 + 0.1\mu g pMBW 96$	None		3603	2785	10349	3463
41	0.3µg pMBW 97 + 0.3µg pMBW 96	None		4903	1054	3453	2380
42	$0.3\mu g pMBW 97 + 0.7\mu g pMBW 96$	euou		578	46	2883	1899
43	None	$0.3\mu$ g GFP	Control	4780	889	25175	8289.6

Table 3. Summary of GFP expression in into CHO cells.transformed by the different experimental constructs.

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## SEQUENCE LISTING

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## We claim :

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- 1) A method for down regulating the expression of a target gene in cells of a eukaryotic organisms, comprising the steps of
  - a) providing the cells of the eukaryotic organism with a chimeric RNA molecule wherein said chimeric RNA molecule comprises
    - a target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene; operably linked to
    - ii) a largely double stranded RNA region; and
  - identifying those eukaryotic organisms wherein the expression of the target gene is down regulated.
- 2) The method according to claim 1, wherein the largely double stranded RNA region comprises a nuclear localization signal from a viroid of the Potato spindle tuber viroid (PSTVd)-type.
- 3) The method according to claim 2, wherein said nuclear localization signal is from a viroid selected from the group consisting of Potato Spindle tuber viroid, Citrus viroid species III, Citrus viroid species IV, Hop latent viroid, Australian grapevine viroid, Tomato planta macho viroid, Coconut tinangaja viroid, Tomato apical stunt viroid, Coconut cadang-cadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid and Citrus bent leaf viroid.
  - 4) The method according to claim 3, wherein said viroid has a genome nucleotide sequence selected from the group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.

- 5) The method according to any one of claims 2 to 4, wherein said nuclear localization signal is from Potato spindle tuber viroid.
- 6) The method according to any one of claims 2 to 5, wherein said nuclear localization signal is from Potato spindle viroid strain RG1.
- 7) The method according to any one of claims 2 to 6, wherein said nuclear localization signal comprises a nucleotide sequence functioning as a nuclear localization signal selected from the nucleotide sequence of SEQ ID N° 3.
- 8) The method according to claim 2 or 3, wherein said largely double stranded RNA comprises a viroid genome nucleotide sequence selected from the group consisting of the genome nucleotide sequence of Potato Spindle tuber viroid, the genome nucleotide sequence of Citrus viroid species III, the genome nucleotide sequence of Citrus viroid species IV, the genome nucleotide sequence of Hop latent viroid, the genome nucleotide sequence of Australian grapevine viroid, the genome nucleotide sequence of Tomato planta macho viroid, the genome nucleotide sequence of Coconut tinangaja viroid, the genome nucleotide sequence of Tomato apical stunt viroid, the genome nucleotide sequence of Coconut cadang-cadang viroid, the genome nucleotide sequence of Citrus exocortis viroid, the genome nucleotide sequence of Columnea latent viroid, the genome nucleotide sequence of Citrus bent leaf viroid.
- 9) The method according to claim 8, wherein said viroid genome nucleotide sequence is selected from group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.

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- 10) The method according to any one of claims 2 to 9, wherein said largely double stranded RNA region comprises a genomic nucleotide sequence of Potato spindle tuber viroid.
- 11) The method according to claim 10, wherein said viroid genome nucleotide sequence is the genome nucleotide sequence of Potato spindle tuber viroid strain RG1.
- 12) The method of claim 11, wherein said genome nucleotide sequence has the nucleotide sequence of SEQ ID N° 3.
  - 13) The method according to claim 1, wherein said largely double stranded RNA region comprises at least 35 repeats of the trinucleotide CUG.
- 14) The method according to claim 13, wherein said largely double stranded RNA region comprises between 44 and 2000 repeats of the trinucleotide CUG.
- 15)The method according to any one of claims 1 to 14, wherein said RNA molecule comprises multiple target-gene specific regions.
  - 16) The method according to any one of claims 1 to 15, wherein said RNA molecule comprises an intron sequence.
- 17) The method according to claim 16, wherein said intron sequence is selected from the group consisting of the pdk2 intron, the catalase intron from Castor bean, the Delta12 desaturase intron from cotton, the Delta 12 desaturase intron from Arabidopsis, the Ubiquitin intron from maize, the Actin intron from rice, the triose phosphate isomerase intron from Aspergillus and the intron from SV40.

- 18) The method according to any one of claims 1 to 17 wherein said eukaryotic organism is a plant.
- 19) The method according to claim 18, wherein said plant is selected from the group of *Arabidopsis*, alfalfa, barley, bean, corn, cotton, flax, pea, rape, rice, rye, safflower, sorghum, soybean, sunflower, tobacco, wheat, asparagus, beet, broccoli, cabbage, carrot, cauliflower, celery, cucmber, eggplant, lettuce, onion, oilseed rape, pepper, potato, pumpkin, radish, spinach, squash, tomato, zucchini, almond, apple, apricot, banana, blackberry, b lueberry, cacao, cherry, coconut, cranberry, date, grape, grapefruit, guava, kiwi, lemon, lime, mango, melon, nectarine, orange, papaya, passion fruit, peach, peanut, pear, pineapple, pistachio, plum, raspberry, strawberry, tangerine, walnut and watermelon.
- 20) The method according to any one of claims 1 to 17, wherein said eukaryotic organism is a fungus, yeast or mold.
  - 21) The method according to any one of claims 1 to 17, wherein said eukaryotic organism is an animal.

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- 22) The method according to claim 21, wherein said animal is a human, mammal, fish, cattle, goat, pig, sheep, rodent, hamster, mouse, rat, guinea pig, rabbit, primate, nematode, shellfish, prawn, crab, lobster, insect, fruit fly, Coleapteran insect, Dipteran insect, Lepidopteran insect and Homeopteran insect.
- 23) The method according to any one of claims 1 to 22, wherein said chimeric RNA is produced by transcription from a chimeric DNA molecule.
- 24) A chimeric RNA molecule for down-regulating the expression of a target gene in a cell of a eukaryotic organisms, comprising

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- a) a target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of said target gene in said cell of said eukaryotic organism; operably linked to
- b) a largely double stranded RNA region; wherein said chimeric RNA molecule, when provided to cells of said eukaryotic organism down-regulates the expression of said target gene.
- 25) The chimeric RNA molecule according to claim 24, wherein the largely double stranded RNA region comprises a nuclear localization signal from a viroid of the Potato spindle tuber viroid (PSTVd) type.
  - 26) The chimeric RNA molecule according to claim 25, wherein said nuclear localization signal is from a viroid selected from the group consisting of Potato Spindle tuber viroid, Citrus viroid species III, Citrus viroid species IV, Hop latent viroid, Australian grapevine viroid, Tomato planta macho viroid, Coconut tinangaja viroid, Tomato apical stunt viroid, Coconut cadangcadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid and Citrus bent leaf viroid.
    - 27) The chimeric RNA molecule according to claim 25 or 26, wherein said viroid has a genome nucleotide sequence selected from the group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.
    - 28) The chimeric RNA molecule according to any one of claims 25 or 26, wherein said nuclear localization signal is from Potato spindle tuber viroid.

- 29) The chimeric RNA molecule according to any one of claims 25 or 26, wherein said nuclear localization signal is from Potato spindle viroid strain RG1.
- 5 30)The chimeric RNA molecule according to any one of claims 25 to 29, wherein said nuclear localization signal comprises a nucleotide sequence functioning as a nuclear localization signal selected from the nucleotide sequence of SEQ ID N° 3.
- 31) The chimeric RNA molecule according to claim 25 or 26, wherein said 10 largely double stranded RNA comprises a viroid genome nucleotide sequence selected from the group consisting of the genome nucleotide sequence of Potato Spindle tuber viroid, the genome nucleotide sequence of Citrus viroid species III, the genome nucleotide sequence of Citrus viroid species IV, the genome nucleotide sequence of Hop latent viroid, the 15 genome nucleotide sequence of Australian grapevine viroid, the genome nucleotide sequence of Tomato planta macho viroid, the genome nucleotide sequence of Coconut tinangaja viroid, the genome nucleotide sequence of Tomato apical stunt viroid, the genome nucleotide sequence of Coconut cadang-cadang viroid, the genome nucleotide sequence of Citrus exocortis 20 viroid, the genome nucleotide sequence of Columnea latent viroid, the genome nucleotide sequence of Hop stunt viroid and the genome nucleotide sequence of Citrus bent leaf viroid.
- 32) The chimeric RNA molecule according to claim 31, wherein said viroid genome nucleotide sequence is selected from group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.

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- 33) The chimeric RNA molecule according to any one of claims 25 to 32, wherein said largely double stranded RNA region comprises a genomic nucleotide sequence of Potato spindle tuber viroid.
- 5 34) The chimeric RNA molecule according to claim 33, wherein said viroid genome nucleotide sequence is the genome nucleotide sequence of Potato spindle tuber viroid strain RG1.
- 35)The chimeric RNA molecule of claim 34, wherein said genome nucleotide sequence has the nucleotide sequence of SEQ ID N° 3.
  - 36)The chimeric RNA molecule according to claim 24, wherein said largely double stranded RNA region comprises at least 35 repeats of the trinucleotide CUG.
  - 37) The chimeric RNA molecule according to claim 36, wherein said largely double stranded RNA region comprises between 44 and 2000 repeats of the trinucleotide CUG.
- 38) The chimeric RNA molecule according to any one of claims 24 to 37, wherein said RNA molecule comprises multiple target-gene specific regions.
  - 39) The chimeric RNA molecule according to any one of claims 24 to 38, wherein said RNA molecule comprises an intron sequence.
  - 40) The chimeric RNA molecule according to claim 39, wherein said intron sequence is selected from the group consisting of the pdk2 intron, the catalase intron from Castor bean, the Delta12 desaturase intron from cotton, the Delta 12 desaturase intron from *Arabidopsis*, the Ubiquitin intron from maize, the Actin intron from rice, the triose phosphate isomerase intron from Aspergillus and the intron from SV40.

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- 41) A chimeric DNA molecule for reduction of the expression of a target gene in a cell of a eukaryotic organism, comprising
  - a) a promoter or promoter region capable of being recognized by RNA polymerases in said cells of said eukaryotic organism; operably linked to
  - b) a DNA region, which when transcribed yields an RNA molecule, said RNA molecule comprising
    - i) a target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of said target gene in said cell of said eukaryotic organism; operably linked to
  - ii) a largely double stranded RNA region; wherein said chimeric DNA molecule, when provided to cells of said eukaryotic organism reduces the expression of said target gene.
- 42) The chimeric DNA molecule according to claim 41, wherein the largely double stranded RNA region comprises a nuclear localization signal from a viroid of the potato spindle tuber viroid type.
- 43) The chimeric DNA molecule according to claim 42, wherein said nuclear localization signal is from a viroid selected from the group consisting of Potato Spindle tuber viroid, Citrus viroid species III, Citrus viroid species IV, Hop latent viroid, Australian grapevine viroid, Tomato planta macho viroid, Coconut tinangaja viroid, Tomato apical stunt viroid, Coconut cadangcadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid and Citrus bent leaf viroid.
- 30 44) The chimeric DNA molecule according to claim 42 or 43, wherein said viroid has a genome nucleotide sequence selected from the group

consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.

- 45) The chimeric DNA molecule according to any one of claims 42 to 44, wherein said nuclear localization signal is from Potato spindle tuber viroid.
  - 46) The chimeric DNA molecule according to claim 45, wherein said nuclear localization signal is from Potato spindle viroid strain RG1.
- 47) The chimeric DNA molecule according to any one of claims 42 to 46, wherein said nuclear localization signal comprises a nucleotide sequence functioning as a nuclear localization signal selected from the nucleotide sequence of SEQ ID N° 3.
- 48) The chimeric DNA molecule according to claim 42 or 43, wherein said 15 largely double stranded RNA comprises a viroid genome nucleotide sequence selected from the group consisting of the genome nucleotide sequence of Potato Spindle tuber viroid, the genome nucleotide sequence of Citrus viroid species III, the genome nucleotide sequence of Citrus viroid species IV, the genome nucleotide sequence of Hop latent viroid, the 20 genome nucleotide sequence of Australian grapevine viroid, the genome nucleotide sequence of Tomato planta macho viroid, the genome nucleotide sequence of Coconut tinangaja viroid, the genome nucleotide sequence of Tomato apical stunt viroid, the genome nucleotide sequence of Coconut cadang-cadang viroid, the genome nucleotide sequence of Citrus exocortis 25 viroid, the genome nucleotide sequence of Columnea latent viroid, the genome nucleotide sequence of Hop stunt viroid and the genome nucleotide sequence of Citrus bent leaf viroid.
- 49) The chimeric DNA molecule according to claim 48, wherein said viroid genome nucleotide sequence is selected from group consisting of SEQ ID

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- N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.
- 50) The chimeric DNA molecule according to any one of claims 42 to 49, wherein said largely double stranded RNA region comprises a genomic nucleotide sequence of Potato spindle tuber viroid.
  - 51)The chimeric DNA molecule according to claim 50, wherein said viroid genome nucleotide sequence is the genome nucleotide sequence of Potato spindle tuber viroid strain RG1.
  - 52) The chimeric DNA molecule of claim 51, wherein said genome nucleotide sequence has the nucleotide sequence of SEQ ID N° 3.
- 53) The chimeric DNA molecule according to claim 41, wherein said largely double stranded RNA region comprises at least 35 repeats of the trinucleotide CUG.
  - 54) The chimeric DNA molecule according to claim 53, wherein said largely double stranded RNA region comprises between 44 and 2000 repeats of the trinucleotide CUG.
    - 55) The chimeric DNA molecule according to any one of claims 41 to 54, wherein said RNA molecule comprises multiple target-gene specific regions.
    - 56) The chimeric DNA molecule according to any one of claims 41 to 55, wherein said RNA molecule comprises an intron sequence.
- 57)The chimeric DNA molecule according to claim 56, wherein said intron sequence is selected from the group consisting of the pdk2 intron, the catalase intron from Castor bean, the Delta12 desaturase intron from

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cotton, the Delta 12 desaturase intron from *Arabidopsis*, the Ubiquitin intron from maize, the Actin intron from rice, the triose phosphate isomerase intron from Aspergillus and the intron from SV40.

- 5 58) The chimeric DNA molecule according to any one of claims 41 to 56, further comprising a transcription termination and polyadenylation signal operably linked to said DNA region encoding said RNA molecule.
  - 59) The chimeric DNA molecule according to any one of claims 41 to 58, wherein said promoter or promoter region is a plant-expressible promoter.
    - 60) The chimeric DNA molecule according to any one of claims 41 to 58, wherein said promoter or promoter region is a promoter which functions in animals.
    - 61)The chimeric DNA molecule according to any one of claims 41 to 58, wherein said promoter or promoter region is a promoter which functions in yeast, fungi or molds.
- 20 62) The chimeric DNA molecule according to any one of claims 41 to 58, wherein said promoter or promoter region is promoter recognized by a single subunit bacteriophage RNA polymerase.
- 63)A cell from a eukaryotic organism comprising a chimeric DNA molecule according to any one of claims 41 to 62.
  - 64)A eukaryotic cell comprising a chimeric RNA molecule according to any one of claims 24 to 40.
- 30 65)The cell according to claim 63 or claim 64, wherein said eukaryotic organism is a plant.

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- 66) The cell according to claim 65, wherein said plant is selected from the group of *Arabidopsis*, alfalfa, barley, bean, corn, cotton, flax, pea, rape, rice, rye, safflower, sorghum, soybean, sunflower, tobacco, wheat, asparagus, beet, broccoli, cabbage, carrot, cauliflower, celery, cucmber, eggplant, lettuce, onion, oilseed rape, pepper, potato, pumpkin, radish, spinach, squash, tomato, zucchini, almond, apple, apricot, banana, blackberry, b lueberry, cacao, cherry, coconut, cranberry, date, grape, grapefruit, guava, kiwi, lemon, lime, mango, melon, nectarine, orange, papaya, passion fruit, peach, peanut, pear, pineapple, pistachio, plum, raspberry, strawberry, tangerine, walnut and watermelon.
- 67) The cell according to claim 63 or claim 64, wherein said eukaryotic organism is a fungus, yeast or mold.
- 68) The cell according to claim 63 or claim 64, wherein said eukaryotic organism is an animal.
- 69) The cell according to claim 68, wherein said animal is a human, mammal, fish, cattle, goat, pig, sheep, rodent, hamster, mouse, rat, guinea pig, rabbit, primate, nematode, shellfish, prawn, crab, lobster, insect, fruit fly, Coleapteran insect, Dipteran insect, Lepidopteran insect and Homeopteran insect.
- 70) A non-human eukaryotic organism, comprising in its cells a chimeric DNA molecule according to any one of claims 41 to 62.
  - 71)A non-human eukaryotic organism, comprising in its cells a chimeric RNA molecule according to any one of claims 24 to 40.

- 72) The non-human eukaryotic organism according to claim 70 or claim 71, wherein said eukaryotic organism is a plant.
- 73) The non-human eukaryotic organism according to claim 72, wherein said plant is selected from the group of *Arabidopsis*, alfalfa, barley, bean, corn, cotton, flax, pea, rape, rice, rye, safflower, sorghum, soybean, sunflower, tobacco, wheat, asparagus, beet, broccoli, cabbage, carrot, cauliflower, celery, cucmber, eggplant, lettuce, onion, oilseed rape, pepper, potato, pumpkin, radish, spinach, squash, tomato, zucchini, almond, apple, apricot, banana, blackberry, b lueberry, cacao, cherry, coconut, cranberry, date, grape, grapefruit, guava, kiwi, lemon, lime, mango, melon, nectarine, orange, papaya, passion fruit, peach, peanut, pear, pineapple, pistachio, plum, raspberry, strawberry, tangerine, walnut and watermelon.
- 74) The non-human eukaryotic organism according to claim 70 or claim 71, wherein said eukaryotic organism is a fungus, yeast or mold.
  - 75) The non-human eukaryotic organism according to claim 70 or claim 71, wherein said eukaryotic organism is an animal.

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- 76) The non-human eukaryotic organism according to claim 75, wherein said animal is a human, mammal, fish, cattle, goat, pig, sheep, rodent, hamster, mouse, rat, guinea pig, rabbit, primate, nematode, shellfish, prawn, crab, lobster, insect, fruit fly, Coleapteran insect, Dipteran insect, Lepidopteran insect and Homeopteran insect.
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- 77) Use of a chimeric RNA molecule according to any one of claims 24 to 40 for reduction of the expression of a target gene in a cell of a eukaryotic organism.

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- 78) Use of a chimeric DNA molecule according to any one of claims 41 to 62 for reduction of the expression of a target gene in a cell of a eukaryotic organism.
- 5 79)A method for making a transgenic eukaryotic organism wherein expression of a target gene in cells of said organism is reduced, said method comprising the steps of :
  - a) providing a chimeric DNA molecule according to any one of claims 41 to
     62 to a cell or cells of said organism to make a transgenic cell or cells;
- b) growing or regenerating a transgenic eukaryotic organism from said transgenic cell or cells.
  - 80)A method for down regulating the expression of a target gene in cells of a eukaryotic organisms, comprising the steps of
    - a) providing the cells of the eukaryotic organism with a first and second chimeric RNA molecule,
      - said first chimeric RNA molecule comprising an antisense targetgene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene;
      - ii) said second chimeric RNA molecule comprising a sense target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity to the complement of said first chimeric RNA molecule;
      - iii) said first and second chimeric RNA being capable of basepairing at least between said 19 consecutive nucleotides of said first chimeric RNA and said 19 consecutive nucleotides of said second chimeric RNA; and
- 30 iv) wherein either said first or said second chimeric RNA molecule comprises a largely double stranded RNA region operably linked to

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- said antisense target-specific RNA region or to said sense target-specific RNA region; and
- b) identifying those eukaryotic organisms wherein the expression of the target gene is down regulated.
- 81) The method according to claim 80, wherein said first and said second chimeric RNA molecule comprise a largely double stranded RNA region.
- 82) The method according to claim 81, wherein said first and said second chimeric RNA molecule comprise the same largely double stranded RNA region.
- 83) The method according to any one of claims 80 to 82, wherein the largely double stranded RNA region comprises a nuclear localization signal from a viroid of the Potato spindle tuber viroid (PSTVd)-type.
- 84) The method according to claim 83, wherein said nuclear localization signal is from a viroid selected from the group consisting of Potato Spindle tuber viroid, Citrus viroid species III, Citrus viroid species IV, Hop latent viroid, Australian grapevine viroid, Tomato planta macho viroid, Coconut tinangaja viroid, Tomato apical stunt viroid, Coconut cadang-cadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid and Citrus bent leaf viroid.
- 85) The method according to claim 83, wherein said viroid has a genome nucleotide sequence selected from the group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.
- 86)The method according to any one of claims 83 to 85, wherein said nuclear localization signal is from Potato spindle tuber viroid.

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- 87)The method according to any one of claims 83 to 86, wherein said nuclear localization signal is from Potato spindle viroid strain RG1.
- 88)The method according to any one of claims 83 to 87, wherein said nuclear localization signal comprises a nucleotide sequence functioning as a nuclear localization signal selected from the nucleotide sequence of SEQ ID N° 3.

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- 89) The method according to claim 83 or 84, wherein said largely double stranded RNA comprises a viroid genome nucleotide sequence selected from the group consisting of the genome nucleotide sequence of Potato Spindle tuber viroid, the genome nucleotide sequence of Citrus viroid species III, the genome nucleotide sequence of Citrus viroid species IV, the genome nucleotide sequence of Hop latent viroid, the genome nucleotide sequence of Tomato planta macho viroid, the genome nucleotide sequence of Coconut tinangaja viroid, the genome nucleotide sequence of Tomato apical stunt viroid, the genome nucleotide sequence of Coconut cadang-cadang viroid, the genome nucleotide sequence of Citrus exocortis viroid, the genome nucleotide sequence of Columnea latent viroid, the genome nucleotide sequence of Columnea latent viroid, the genome nucleotide sequence of Citrus bent leaf viroid.
- 90)The method according to claim 89, wherein said viroid genome nucleotide sequence is selected from group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.
  - 91) The method according to any one of claims 83 to 90, wherein said largely double stranded RNA region comprises a genomic nucleotide sequence of Potato spindle tuber viroid.

- 92) The method according to claim 91, wherein said viroid genome nucleotide sequence is the genome nucleotide sequence of Potato spindle tuber viroid strain RG1.
- 5 93)The method of claim 92, wherein said genome nucleotide sequence has the nucleotide sequence of SEQ ID N° 3.
  - 94) The method according to any one of claims 80 to 82, wherein said largely double stranded RNA region comprises at least 35 repeats of the trinucleotide CUG.

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- 95)The method according to claim 94, wherein said largely double stranded RNA region comprises between 44 and 2000 repeats of the trinucleotide CUG.
- 96)The method according to any one of claims 80 to 95, wherein said RNA molecule comprises multiple target-gene specific regions.
- 97)The method according to any one of claims 80 to 96, wherein said RNA molecule comprises an intron sequence.
  - 98) The method according to claim 97, wherein said intron sequence is selected from the group consisting of the pdk2 intron, the catalase intron from Castor bean, the Delta12 desaturase intron from cotton, the Delta 12 desaturase intron from Arabidopsis, the Ubiquitin intron from maize, the Actin intron from rice, the triose phosphate isomerase intron from Aspergillus and the intron from SV40.
  - 99) The method according to any one of claims 80 to 98, wherein said first and second chimeric RNA are transcribed from a first and second chimeric gene.

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- 100) A cell from a eukaryotic organism comprising a first and second chimeric RNA molecule,
  - said first chimeric RNA molecule comprising an antisense targetgene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity with the complement of 19 consecutive nucleotides from the nucleotide sequence of the target gene;
  - ii) said second chimeric RNA molecule comprising a sense target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity to the complement of said first chimeric RNA molecule;
  - iii) said first and second chimeric RNA being capable of basepairing at least between said 19 consecutive nucleotides of said first chimeric RNA and said 19 consecutive nucleotides of said second chimeric RNA; and
  - iv) wherein either said first or said second chimeric RNA molecule comprises a largely double stranded RNA region operably linked to said antisense target-specific RNA region or to said sense targetspecific RNA region.
- 101) The cell according to claim 100, wherein said first and said second chimeric RNA molecule comprise a largely double stranded RNA region.
- 25 102) The cell according to claim 101, wherein said first and said second chimeric RNA molecule comprise the same largely double stranded RNA region.
- 103) The cell according to any one of claims 100 to 102, wherein the largely double stranded RNA region comprises a nuclear localization signal from a viroid of the Potato spindle tuber viroid (PSTVd)-type.

- 104) The cell according to claim 103, wherein said nuclear localization signal is from a viroid selected from the group consisting of Potato Spindle tuber viroid, Citrus viroid species III, Citrus viroid species IV, Hop latent viroid, Australian grapevine viroid, Tomato planta macho viroid, Coconut tinangaja viroid, Tomato apical stunt viroid, Coconut cadang-cadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid and Citrus bent leaf viroid.
- 10 105) The cell according to claim 103, wherein said viroid has a genome nucleotide sequence selected from the group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.
  - 106) The cell according to any one of claims 103 to 105, wherein said nuclear localization signal is from Potato spindle tuber viroid.
    - 107) The cell according to any one of claims 103 to 106, wherein said nuclear localization signal is from Potato spindle viroid strain RG1.
- 20 108) The cell according to any one of claims 103 to 107, wherein said nuclear localization signal comprises a nucleotide sequence functioning as a nuclear localization signal selected from the nucleotide sequence of SEQ ID N° 3.
- 25 109) The cell according to claim 103 or 104, wherein said largely double stranded RNA comprises a viroid genome nucleotide sequence selected from the group consisting of the genome nucleotide sequence of Potato Spindle tuber viroid, the genome nucleotide sequence of Citrus viroid species III, the genome nucleotide sequence of Citrus viroid species IV, the genome nucleotide sequence of Hop latent viroid, the genome nucleotide sequence of Australian grapevine viroid, the genome nucleotide sequence

of Tomato planta macho viroid, the genome nucleotide sequence of Coconut tinangaja viroid, the genome nucleotide sequence of Tomato apical stunt viroid, the genome nucleotide sequence of Coconut cadang-cadang viroid, the genome nucleotide sequence of Citrus exocortis viroid, the genome nucleotide sequence of Columnea latent viroid, the genome nucleotide sequence of Hop stunt viroid and the genome nucleotide sequence of Citrus bent leaf viroid.

- 110) The cell according to claim 109, wherein said viroid genome nucleotide sequence is selected from group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.
  - 111) The cell according to any one of claims 103 to 109, wherein said largely double stranded RNA region comprises a genomic nucleotide sequence of Potato spindle tuber viroid.
  - 112) The cell according to claim 111, wherein said viroid genome nucleotide sequence is the genome nucleotide sequence of Potato spindle tuber viroid strain RG1.

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- 113) The cell of claim 112, wherein said genome nucleotide sequence has the nucleotide sequence of SEQ ID N° 3.
- 114) The cell according to any one of claims 100 to 102, wherein said largely double stranded RNA region comprises at least 35 repeats of the trinucleotide CUG.
  - 115) The cell according to claim 114, wherein said largely double stranded RNA region comprises between 44 and 2000 repeats of the trinucleotide CUG.

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- 116) The cell according to any one of claims 100 to 115, wherein said RNA molecule comprises multiple target-gene specific regions.
- 117) The cell according to any one of claims 100 to 116, wherein said RNA molecule comprises an intron sequence.
- 118) The cell according to claim 117, wherein said intron sequence is selected from the group consisting of the pdk2 intron, the catalase intron from Castor bean, the Delta12 desaturase intron from cotton, the Delta 12 desaturase intron from *Arabidopsis*, the Ubiquitin intron from maize, the Actin intron from rice, the triose phosphate isomerase intron from Aspergillus and the intron from SV40.
- 119) The cell according to any one of claims 100 to 118 wherein said first and second chimeric RNA are transcribed from a first and second chimeric gene.
- 120) A non-human eukaryotic organism comprising the cell according to any one of claims 100 to claim 119.
- 121) A chimeric sense RNA molecule for reduction of expression of a target gene in a cell of a eukaryotic organism in cooperation with a chimeric antisense RNA molecule, said chimeric sense RNA molecule comprising
  - a) a sense target-gene specific RNA region comprising a nucleotide sequence of at least about 19 consecutive nucleotides having at least about 94% sequene identity to the nucleotide of said target gene; operably linked to
  - b) a largely double stranded RNA region.

- 122) The chimeric RNA molecule according to claim 121, wherein the largely double stranded RNA region comprises a nuclear localization signal from a viroid of the Potato spindle tuber viroid (PSTVd)-type.
- 5 123) The chimeric RNA molecule according to claim 122, wherein said nuclear localization signal is from a viroid selected from the group consisting of Potato Spindle tuber viroid, Citrus viroid species III, Citrus viroid species IV, Hop latent viroid, Australian grapevine viroid, Tomato planta macho viroid, Coconut tinangaja viroid, Tomato apical stunt viroid, Coconut cadang-cadang viroid, Citrus exocortis viroid, Columnea latent viroid, Hop stunt viroid and Citrus bent leaf viroid.
  - 124) The chimeric RNA molecule according to claim 123, wherein said viroid has a genome nucleotide sequence selected from the group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.
  - 125) The chimeric RNA molecule according to any one of claims 122 to 124, wherein said nuclear localization signal is from Potato spindle tuber viroid.
  - 126) The chimeric RNA molecule according to any one of claims 122 to 125, wherein said nuclear localization signal is from Potato spindle viroid strain RG1.
- 127) The chimeric RNA molecule according to any one of claims 122 to 126, wherein said nuclear localization signal comprises a nucleotide sequence functioning as a nuclear localization signal selected from the nucleotide sequence of SEQ ID N° 3.
- 128) The chimeric RNA molecule according to claim 122 or 123, wherein said largely double stranded RNA comprises a viroid genome nucleotide

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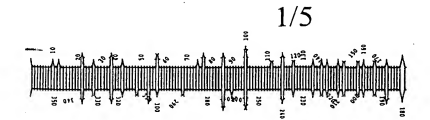
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sequence selected from the group consisting of the genome nucleotide sequence of Potato Spindle tuber viroid, the genome nucleotide sequence of Citrus viroid species III, the genome nucleotide sequence of Citrus viroid species IV, the genome nucleotide sequence of Hop latent viroid, the genome nucleotide sequence of Australian grapevine viroid, the genome nucleotide sequence of Tomato planta macho viroid, the genome nucleotide sequence of Coconut tinangaja viroid, the genome nucleotide sequence of Tomato apical stunt viroid, the genome nucleotide sequence of Coconut cadang-cadang viroid, the genome nucleotide sequence of Citrus exocortis viroid, the genome nucleotide sequence of Columnea latent viroid, the genome nucleotide sequence of Columnea latent viroid, the genome nucleotide sequence of Citrus bent leaf viroid.

- 129) The chimeric RNA molecule according to claim 128, wherein said viroid genome nucleotide sequence is selected from group consisting of SEQ ID N° 3, SEQ ID N° 4, SEQ ID N° 5, SEQ ID N° 6, SEQ ID N° 7 and SEQ ID N° 8.
- 130) The chimeric RNA molecule according to any one of claims 122 to 129, wherein said largely double stranded RNA region comprises a genomic nucleotide sequence of Potato spindle tuber viroid.
  - 131) The chimeric RNA molecule according to claim 130, wherein said viroid genome nucleotide sequence is the genome nucleotide sequence of Potato spindle tuber viroid strain RG1.
  - 132) The chimeric RNA molecule of claim 131, wherein said genome nucleotide sequence has the nucleotide sequence of SEQ ID N° 3.

- 133) The chimeric RNA molecule according to claim 121, wherein said largely double stranded RNA region comprises at least 35 repeats of the trinucleotide CUG.
- 5 134) The chimeric RNA molecule according to claim 133, wherein said largely double stranded RNA region comprises between 44 and 2000 repeats of the trinucleotide CUG.
- 135) The chimeric RNA molecule according to any one of claims 121 to 134, wherein said RNA molecule comprises multiple target-gene specific regions.
  - 136) The chimeric RNA molecule according to any one of claims 121 to 135, wherein said RNA molecule comprises an intron sequence.
- 137) The chimeric RNA molecule according to claim 136, wherein said intron sequence is selected from the group consisting of the pdk2 intron, the catalase intron from Castor bean, the Delta12 desaturase intron from cotton, the Delta 12 desaturase intron from *Arabidopsis*, the Ubiquitin intron from maize, the Actin intron from rice, the triose phosphate isomerase intron from Aspergillus and the intron from SV40.
  - 138) A chimeric DNA molecule for reduction of the expression of a target gene in a cell of a eukaryotic organism, comprising

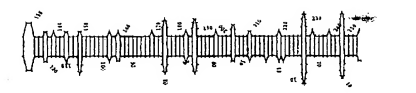
- a) a promoter or promoter region capable of being recognized by RNA polymerases in said cells of said eukaryotic organism; operably linked
- b) a DNA region, which when transcribed yields a chimeric sense RNA molecule as described in any one of claim 121 to 137.



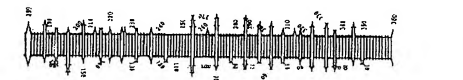
Α



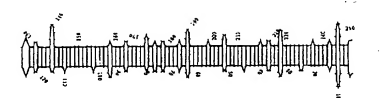
В



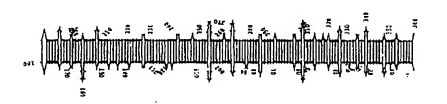
C



D



E



F

Figure 1

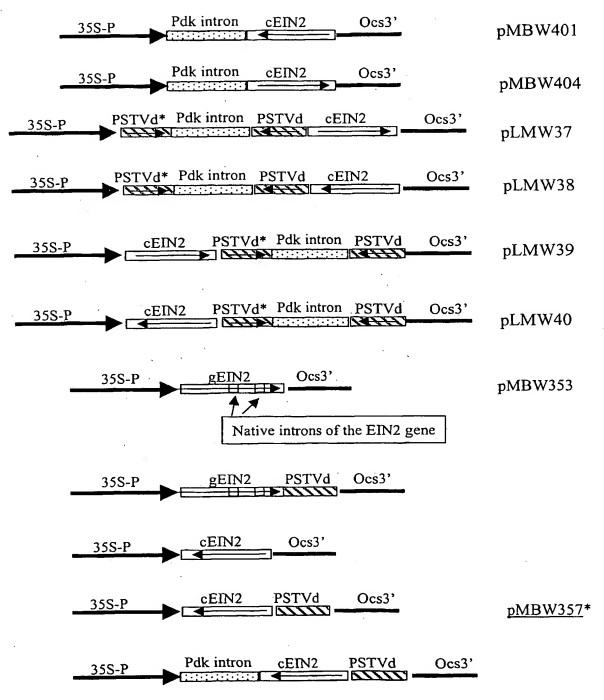
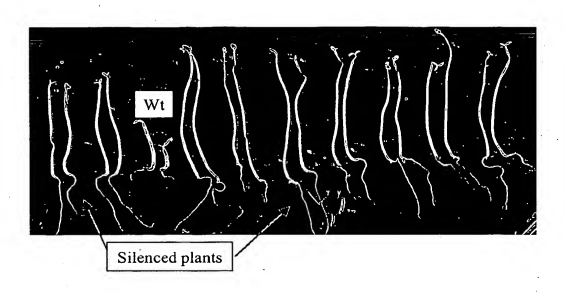


Figure 2

A.



B.

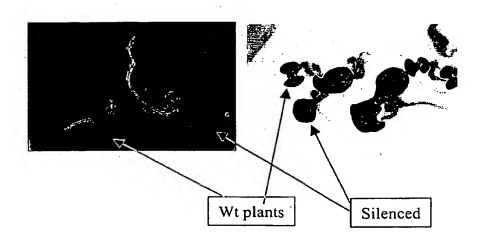


Figure 3

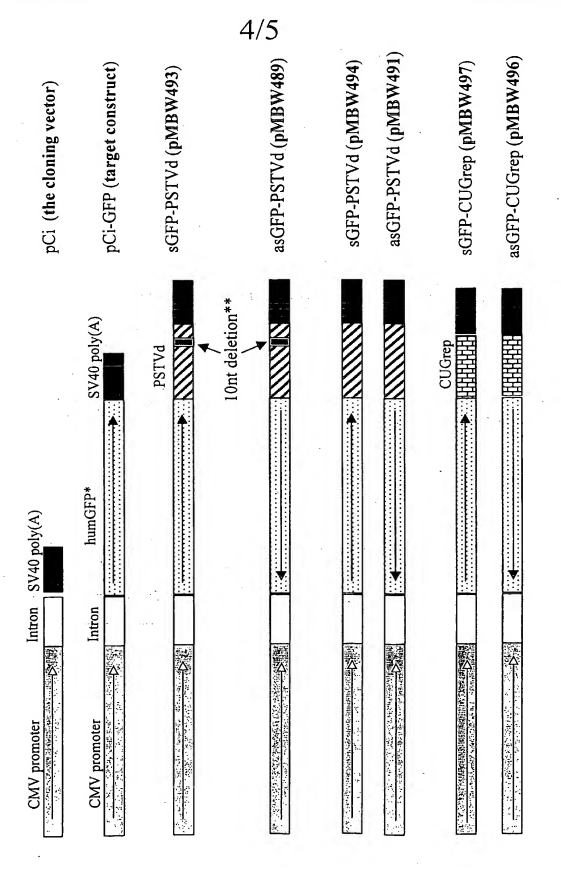


Figure 4

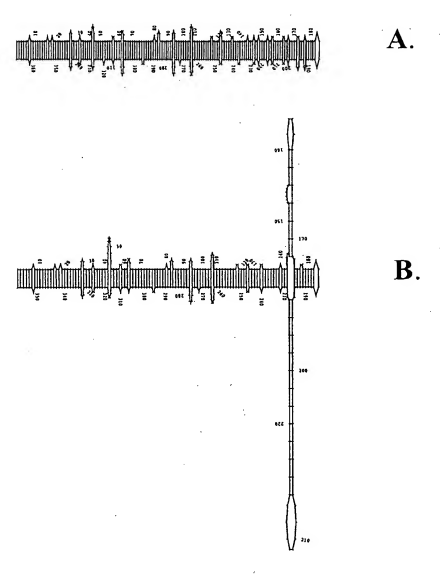


Figure 5

International application No.

PCT/AU03/00292

Α.	CLASSIFICATION OF SUBJECT MATTER					
Int. Cl. 7: C12N 15/11						
According to International Patent Classification (IPC) or to both national classification and IPC						
В.	FIELDS SEARCHED					
Minimum documentation searched (classification system followed by classification symbols) IPC (WPIDS) AND CHEMICAL ABSTRACTS						
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched SEE BELOW						
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) WPIDS, CA, MEDLINE						
C. DOCUMENTS CONSIDERED TO BE RELEVANT						
Category*	Citation of document, with indication, where appr	opriate, of the relevant passages	Relevant to claim No.			
Y	The Plant Journal (2001) 27(6), Wesley et al and high-throughput gene silencing in plants		2-14, 25-37, 42-54, 83-95, 103-115, 122- 134			
Υ	Journal of General Virology (2001) 82, Zhao virus X in a whole plant assay to demonstrate viroid", pages 1491-7		2-12, 25-35, 42-52, 83-93, 103-113, 122- 132			
Y	Proc. Natl. Acad. Sci. USA (1997) 94, Davis repeat in the 3' untranslated region of myotor results in nuclear retention of transcripts", pa	13, 14, 36, 37, 53, 54, 94, 95, 114, 115, 133, 134				
X Further documents are listed in the continuation of Box C See patent family annex						
"A" docume which i relevan	which is not considered to be of particular and not in conflict with the application but cited to understand the principl or theory underlying the invention					
after the	international filing date considered novel or cannot be considered to involve an inventive step when the document is taken alone					
claim(s) publica	nt which may throw doubts on priority or which is cited to establish the cion date of another citation or other special  "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to					
"O" docume	as specified) a person skilled in the art nt referring to an oral disclosure, use, "&" document member of the same patent family on or other means					
"P" docume	on or other means int published prior to the international filing later than the priority date claimed					
Date of the actual completion of the international search  Date of mailing of the international search report 5 - MAY 2003						
23 April 2003						
Name and mailing address of the ISA/AU		Authorized officer				
AUSTRALIAN PATENT OFFICE PO BOX 200, WODEN ACT 2606, AUSTRALIA E-mail address: pct@ipaustralia.gov.au Facsimile No. (02) 6285 3929		CHRISTOPHER LUTON Telephone No: (02) 6283 2256				
	<del></del>					

International application No.
PCT/AU03/00292

Category*	Citation of document, with indication, where appropriate, of the relevant passages		
A	Nucleic Acids Research (2001) 29(11), Papaefthimiou et al., "Replicating potato spindle tuber viroid RNA is accompanied by short RNA fragments that are characteristic of post-transcriptional gene silencing", pages 2395-2400  Molecular Plant-Microbe Interactions (2001) 14(11), Itaya et al., "Potato spindle tuber viroid as Inducer of RNA Silencing in Infected Tomato", pages 1332-4  2-14, 2.  42-54, 8 103-115 134		
Α			
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International application No.

PCT/AU03/00292

Box I	Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)			
This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:				
1.	Claims Nos:			
	because they relate to subject matter not required to be searched by this Authority, namely:			
2.	X Claims Nos: 1, 15-24, 38-41, 55-82, 96-102, 116-121, 135-138			
	because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:  See supplemental box			
3.	Claims Nos:			
	because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a)			
Box II	Observations where unity of invention is lacking (Continuation of item 3 of first sheet)			
This Inte	mational Searching Authority found multiple inventions in this international application, as follows:			
•	A. Carrier and Car			
,				
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims			
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.			
3.	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:			
	·			
4.	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:			
Remark on Protest				
	No protest accompanied the payment of additional search fees.			

International application No.

PCT/AU03/00292

Supp	lemental	Box
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(To be used when the space in any of Boxes I to VIII is not sufficient)

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#### Continuation of Box No: I

Claims 1, 15-24, 38-41, 55-82, 96-102, 116-121, 135-138 do not define the matter for which protection is sought in terms of the technical features of the invention (see Rule 6.3(a), Part B: Rules Concerning Chapter I of the Treaty). The specification, when read as a whole, indicates that the invention relates to the use of either viroid sequences or trinucleotide repeat sequences to target the antisense RNA to the nucleus. Claims 1, 15-24, 38-41, 55-82, 96-102, 116-121 and 135-138 are not limited to the use of such sequences and therefore do not define the subject matter for which protection is sought in terms of the technical features of the invention.